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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

WO 88/00205 (51) International Patent Classification 4: (11) International Publication Number: C07K 13/00, 15/00, A61K 37/00 (43) International Publication Date: 14 January 1988 (14.01.88) C12P 21/00, 21/02, C12N 15/00 C07H 15/12 Boston, MA 02116 (US). PCT/US87/01537 (21) International Application Number: 30 June 1987 (30.06.87) (22) International Filing Date: (31) Priority Application Numbers: 880,776 943,332 028,285

1 July 1986 (01.07.86) (32) Priority Dates: 17 December 1986 (17.12.86) 20 March 1987 (20.03.87)

26 March 1987 (26.03.87)

(33) Priority Country:

031,346

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(81) Designated States: AT (European patent), AU, BE (European patent), BG, BJ (OAPI patent), CF (OAPI patent), CG (OAPI patent), CH (European patent), CM (OAPI patent), DE (European patent), DK, FI, FR (European patent), GA (OAPI patent), GB, GB (European patent), HU, IT (European patent), JP, KR, (OAPI patent), JP, KB, (OAPI paten LU (European patent), ML (OAPI patent), MR (OA-PI patent), NL (European patent), NO, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent).

Published

With international search report. With amended claims.

(54) Title: NOVEL OSTEOINDUCTIVE COMPOSITIONS

(57) Abstract

Human and bovine bone inductive factor products and processes. The factors may be produced by recombinant techniques and are useful in the research and treatment of bone and periodontal defects.

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NOVEL OSTEOINDUCTIVE COMPOSITIONS

The present invention relates to novel proteins and processes for obtaining them. These proteins are capable of inducing cartilage and bone formation.

Background

Bone is a highly specialized tissue characterized by an extensive matrix structure formed of fibrous bundles of the protein collagen, and proteoglycans, noncollagenous proteins, lipids and acidic proteins. The processes of bone formation and renewal/repair of bone tissue, which occur continuously throughout life, are performed by specialized cells. Normal embryonic long bone development is preceded by formation of a cartilage model. Bone growth is presumably mediated by "osteoblasts" (bone-forming cells), while remodeling of bone is apparently accomplished by the joint activities of bone-resorbing cells, called "osteoclasts" and osteoblasts. A variety of osteogenic, cartilage-inducing and bone inducing factors have been described. See, e.g. European patent applications 148,155 and 169,016 for discussions thereof.

Brief Description of the Invention

The present invention provides novel proteins in purified form. Specifically, four of the novel proteins are designated BMP-1, BMP-2 Class I (or BMP-2), BMP-3, and BMP-2 Class II (or BMP-4) wherein BMP is bone morphogenic protein. These proteins are characterized by peptide sequences the same as or substantially homologous to amino acid sequences illustrated in Tables II through VIII below. They are capable of inducing bone formation at a predetermined site. These bone inductive factors are further characterized by biochemical and biological characteristics including activity at a concentration of 10 to 1000ng/gram of bone in an in vivo rat bone formation assay described below. Proteins of this invention may be encoded by the DNA sequences depicted in the Tables or by sequences capable

of hybridizing thereto and coding for polypeptides with bone growth factor biological properties or other variously modified sequences demonstrating such properties.

One of the proteins of the invention is designated BMP-A portion of the human BMP-1 or hBMP-1 is characterized by the same or substantially the same peptide sequence as that of amino acid #1 through amino acid #37 of Table V, below which represents a genomic hBMP-1 fragment or amino acid #1 through amino acid #730 of Table VI which represents the hBMP-1 cDNA. hBMP-1 or a related bone inductive factor may be further characterized by at least a portion of these These peptide sequences are encoded by the same or: substantially the same DNA sequence, as depicted in nucleotide #3440 through nucleotide #3550 of Table V and in nucleotide #36 through nucleotide #2225 of Table VI, respectively. These hBMP-1 polypeptides are further characterized by the ability to induce bone formation. 1 demonstrates activity in an in vivo rat bone formation assay at a concentration of 10 to 1000ng/gram of bone.

The homologous bovine growth factor of the invention, designated bBMP-1, is characterized by a peptide sequence containing the same or substantially the same sequence as that of amino acid #1 through amino acid #37 of Table II below which represents a genomic bBMP-1 fragment. This peptide sequence is encoded by the same or substantially the same DNA sequence as depicted in nucleotide #294 through nucleotide #404 of Table II. The bovine peptide sequence identified in Table II below is also 37 amino acids in length. bBMP-1 is further characterized by the ability to induce bone formation.

Another bone inductive protein composition of the invention is designated BMP-2 Class I (or BMP-2). It is characterized by at least a portion of a peptide sequence the same or substantially the same as that of amino acid #1 through amino acid #396 of Table VII which represents the cDNA hBMP-2 Class I. This peptide sequence is encoded by the same or

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substantially the same DNA sequence, as depicted in nucleotide #356 through nucleotide #1543 of Table VII. The human peptide sequence identified in Table VII is 396 amino acids in length. hBMP-2 or related bone inductive proteins may also be characterized by at least a portion of this peptide sequence. hBMP-2 Class I is further characterized by the ability to induce bone formation.

The homologous bovine bone inductive protein of the invention designated bBMP-2 Class I (or bBMP-2), has a DNA sequence identified in Table III below which represents the genomic sequence. This bovine DNA sequence has a prospective 129 amino acid coding sequence followed by approximately 205 nucleotides (a presumptive 3' non-coding sequence). bBMP-2, Class I is further characterized by the ability to induce bone formation. A further bone inductive protein composition of the invention is designated BMP-2 Class II or BMP-4. human protein hBMP-2 Class II (or hBMP-4) is characterized by at least a portion of the same or substantially the same peptide sequence between amino acid #1 through amino acid #408 of Table VIII, which represents the cDNA of hBMP-2 Class II. peptide sequence is encoded by at least a portion of the same or substantially the same DNA sequence as depicted in nucleotide #403 through nucleotide #1626 of Table VIII. This factor is further characterized by the ability to induce bone formation.

Still another bone inductive factor of the invention, BMP-3, is represented by the bovine homolog bBMP-3. bBMP-3 is characterized by the DNA sequence and amino acid sequence of Table IV A and B which represents the bovine genomic sequence. It is characterized by at least a portion of a peptide sequence the same or substantially the same as amino acid #1 through amino acid #175 of Table IV A and B. BMP-3 is further characterized by the ability to induce bone formation. The bovine factor may be employed as a tool for obtaining the analogous human BMP-3 protein or other mammalian bone inductive proteins. The proper characterization of this bovine bone

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inductive factor provides the essential "starting point" for the method employing this sequence. The method, employing techniques known to those skilled in the art of genetic engineering, involves using the bovine DNA sequence as a probe to screen a human genomic or cDNA library; and identifying the DNA sequences which hybridize to the probes. A clone with a hybridizable sequence is plaque purified and the DNA isolated therefrom, subcloned and subjected to DNA sequence analysis. Thus as another aspect of this invention is a human protein hBMP-3, produced by this method.

Another aspect of the invention provides pharmaceutical compositions containing a therapeutically effective amount of one or more bone growth factor polypeptides according to the invention in a pharmaceutically acceptable vehicle. These compositions may further include other therapeutically useful agents. They may also include an appropriate matrix for delivering the proteins to the site of the bone defect and for providing a structure for bone growth. These compositions may be employed in methods for treating a number of bone defects and periodontal disease. These methods, according to the invention, entail administering to a patient needing such bone formation an effective amount of at least one of the novel proteins BMP-1, BMP-2 Class I, BMP-2 Class-II, and BMP-3 as described herein.

Still a further aspect of the invention are DNA sequences coding on expression for a human or bovine polypeptide having the ability to induce bone formation. Such sequences include the sequence of nucleotides in a 5' to 3' direction illustrated in Tables II through VIII. Alternatively, a DNA sequence which hybridizes under stringent conditions with the DNA sequences of Tables II - VIII or a DNA sequence which hybridizes under non-stringent conditions with the illustrated DNA sequences and which codes on expression for a protein having at least one bone growth factor biological property are included in the present invention. Finally, allelic or other variations of the

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sequences of Tables II through VIII, whether such nucleotide changes result in changes in the peptide sequence or not, are also included in the present invention.

Still a further aspect of the invention is a vector containing a DNA sequence as described above in operative association with an expression control sequence. Such vector may be employed in a novel process for producing a bone growth factor polypeptide in which a cell line transformed with a DNA sequence encoding expression of a bone growth factor polypeptide in operative association with an expression control sequence therefor, is cultured. This claimed process may employ a number of known cells as host cells for expression of the polypeptide. Presently preferred cell lines are mammalian cell lines and bacterial cells.

Other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description and preferred embodiments thereof.

Detailed Description of the Invention

The proteins of the present invention are characterized by amino acid sequences or portions thereof the same as or substantially homologous to the sequences shown in Tables II - VIII below. These proteins are also characterized by the ability to induce bone formation.

The bone growth factors provided herein also include factors encoded by the sequences similar to those of Tables II - VIII, but into which modifications are naturally provided (e.g. allelic variations in the nucleotide sequence which may result in amino acid changes in the polypeptide) or deliberately engineered. For example, synthetic polypeptides may wholly or partially duplicate continuous sequences of the amino acid residues of Tables II - VIII. These sequences, by virtue of sharing primary, secondary, or tertiary structural and conformational characteristics with bone growth factor polypeptides of Tables II - VIII may possess bone growth factor biological properties in common therewith. Thus, they may be

employed as biologically active substitutes for naturallyoccurring bone growth factor polypeptides in therapeutic processes.

Other specific mutations of the sequences of the bone growth factors described herein involve modifications of one or both of the glycosylation sites. The absence of glycosylation or only partial glycosylation results from amino acid substitution or deletion at one or both of the asparagine-linked glycosylation recognition sites present in the sequences of the bone growth factors shown in Tables II-VIII. The asparagine-linked glycosylation recognition sites comprise tripeptide sequences which are specifically recognized by appropriate cellular glycosylation enzymes. These tripeptide sequences are either asparagine-X-threonine or asparagine-Xserine, where X is usually any amino acid. A variety of amino acid substitutions or deletions at one or both of the first or third amino acid positions of a glycosylation recognition site (and/or amino acid deletion at the second position) results in non-glycosylation at the modified tripeptide sequence.

The present invention also encompasses the novel DNA sequences, free of association with DNA sequences encoding other proteinaceous materials, and coding on expression for bone growth factors. These DNA sequences include those depicted in Tables II - VIII in a 5' to 3' direction and those sequences which hybridize under stringent hybridization conditions [see, T. Maniatis et al, Molecular Cloning (A Laboratory Manual), Cold Spring Harbor Laboratory (1982), pages 387 to 389] to the DNA sequences of Tables II - VIII.

DNA sequences which hybridize to the sequences of Tables II - VIII under relaxed hybridization conditions and which code on expression for bone growth factors having bone growth factor biological properties also encode bone growth factors of the invention. For example, a DNA sequence which shares regions of significant homology, e.g., sites of glycosylation

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or disulfide linkages, with the sequences of Tables II - VIII and encodes a bone growth factor having one or more bone growth factor biological properties clearly encodes a member of this novel family of growth factors, even if such a DNA sequence would not stringently hybridize to the sequence of Tables II - VIII.

Similarly, DNA sequences which code for bone growth factor polypeptides coded for by the sequences of Tables II - VIII, but which differ in codon sequence due to the degeneracies of the genetic code or allelic variations (naturally-occurring base changes in the species population which may or may not result in an amino acid change) also encode the novel growth factors described herein. Variations in the DNA sequences of Tables II -VIII which are caused by point mutations or by induced modifications to enhance the activity, half-life or production of the polypeptides encoded thereby are also encompassed in the invention.

Another aspect of the present invention provides a novel method for producing the novel osteoinductive factors. method of the present invention involves culturing a suitable cell or cell line, which has been transformed with a DNA sequence coding on expression for a novel bone growth factor polypeptide of the invention, under the control of known Suitable cells or cell lines may be regulatory sequences. mammalian cells, such as Chinese hamster ovary cells (CHO). The selection of suitable mammalian host cells and methods for transformation, culture, amplification, screening and product production and purification are known in the art. See, e.g., Gething and Sambrook, Nature, 293:620-625 (1981), or alternatively, Kaufman et al, Mol. Cell. Biol., 5(7):1750-1759 (1985) or Howley et al, U.S. Patent 4,419,446. Another suitable mammalian cell line, which is described in the accompanying examples, is the monkey COS-1 cell line. A similarly useful mammalian cell line is the CV-1 cell line.

Bacterial cells are suitable hosts. For example, the

various strains of <u>E</u>. <u>coli</u> (e.g., HB101, MC1061) are well-known as host cells in the field of biotechnology. Various strains of <u>B</u>: <u>subtilis</u>, <u>Pseudomonas</u>, other bacilli and the like may also be employed in this method.

Many strains of yeast cells known to those skilled in the art are also available as host cells for expression of the polypeptides of the present invention. Additionally, where desired, insect cells may be utilized as host cells in the method of the present invention. See, e.g. Miller et al, Genetic Engineering, 8:277-298 (Plenum Press 1986) and references cited therein.

Another aspect of the present invention provides vectors for use in the method of expression of these novel osteoinductive polypeptides. Preferably the vectors contain the full novel DNA sequences described above which code for the novel factors of the invention. Additionally the vectors also contain appropriate expression control sequences permitting expression of the bone inductive protein sequences. Alternatively, vectors incorporating modified sequences as described above are also embodiments of the present invention and useful in the production of the bone inductive proteins. The vectors may be employed in the method of transforming cell lines and contain selected regulatory sequences in operative association with the DNA coding sequences of the invention which are capable of directing the replication and expression thereof in selected host cells. Useful regulatory sequences for such vectors are known to one of skill in the art and may be selected depending upon the selected host Such selection is routine and does not form part of the present invention.

A protein of the present invention, which induces bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures. An osteogenic preparation employing one or more of the proteins of the invention may have prophylactic use in closed as well as open

fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery. An osteogenic factor of the invention may be valuable in the treatment of periodontal disease, and in other tooth repair processes. Such agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of progenitors of bone-forming cells. Of course, the proteins of the invention may have other therapeutic uses.

A further aspect of the invention is a therapeutic method and composition for repairing fractures and other conditions related to bone defects or periodontal diseases. composition comprises a therapeutically effective amount of at least one of the bone inductive factor proteins of the The bone inductive factors according to the invention. present invention may be present in a therapeutic composition in admixture with a pharmaceutically acceptable vehicle or matrix. Further therapeutic methods and compositions of the invention comprise a therapeutic amount of a bone inductive factor of the invention with a therapeutic amount of at least one of the other bone inductive factors of the invention. Additionally, the proteins according to the present invention or a combination of the proteins of the present invention may be co-administered with one or more different osteoinductive factors with which it may interact. Further, the bone inductive proteins may be combined with other agents beneficial to the treatment of the bone defect in question. Such agents include, but are not limited to various growth factors. The preparation of such physiologically acceptable protein compositions, having due regard to pH, isotonicity, stability and the like, is within the skill of the art.

In particular, BMP-1 may be used individually in a

composition. BMP-1 may also be used in combination with one or more of the other proteins of the invention. BMP-1 and BMP-2 Class I may be used in combination. BMP-1 and BMP-3 may be used in combination. Furthermore, BMP-1 may be used in combination with two or three of the other proteins of the invention. For example, BMP-1, BMP-2 Class I, and BMP-2 Class II may be combined. BMP-1 may also be combined with BMP-2 Class I, and BMP-3. Further, BMP-1 may be combined with BMP-2 Class II, and BMP-3. BMP-1, BMP-2 Class I, BMP-2 Class II, and BMP-3. BMP-1, BMP-2 Class I, BMP-2 Class II, and BMP-3 may be combined.

BMP-2 Class I may be used individually in a pharmaceutical composition. BMP-2 Class I may also be used in combination with one or more of the other proteins of the invention. BMP-2 Class I may be combined with BMP-2 Class II. It may also be combined with BMP-3. Further BMP-2 Class I may be combined with BMP-2 Class II and BMP-3.

BMP-2 Class II may be used individually in pharmaceutical composition. In addition, it may be used in combination with other proteins as identified above. Further it may be used in combination with BMP-3.

BMP-3 may be used individually in a composition. It may further be used in the various combinations identified above.

The therapeutic method includes locally administering the composition as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition may desirably be encapsulated or injected in a viscous form for delivery to the site of bone damage. Preferably, the bone growth inductive factor composition would include a matrix capable of delivering the bone inductive factor to the site of bone damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices may be formed of other materials presently in use for other implanted medical

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applications.

The choice of material is based on, for example, biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. Similarly, the application of the osteoinductive factors will define the Potential matrices for the appropriate formulation. osteoinductive factors may be biodegradable and chemically defined, such as, but not limited to calcium sulfate, tricalciumphosphate, hydroxyapatite, polylactic acid, polyanhydrides; biodegradable and biologically well defined, such as bone or dermal collagen, other pure proteins or extracellular matrix components; nonbiodegradable and chemically defined, such as sintered hydroxyapatite, bioglass, aluminates, or other ceramics; or combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalciumphosphate. The bioceramics might also be altered in composition, such as in calcium-aluminatephosphate and processing to alter for example, pore size, particle size, particle shape, and biodegradability.

The dosage regimen will be determined by the attending physician considering various factors which modify the action of such a growth factor, e.g. amount of bone weight desired to be formed, the site of bone damage, the condition of the damaged bone, the patient's age, sex, and diet, the severity of any infection, time of administration and other clinical factors. The dosage may vary with the type of matrix used in the reconstitution and the composition of BMP's. The addition of other known growth factors, such as IGF 1 (insulin like growth factor 1), to the final composition, may also effect the dosage. Generally, the dosage regimen should be in the range of approximately 10 to 106 nanograms of protein per gram of bone weight desired. Progress can be monitored by periodic assessment of bone growth and/or repair, e.g. x-rays. therapeutic compositions are also presently valuable for veterinary applications due to the lack of species specificity

in bone inductive factors. Particularly domestic animals and thoroughbred horses in addition to humans are desired patients for such treatment with the bone inductive factors of the present invention.

The following examples illustrate practice of the present invention in recovering and characterizing the bovine proteins and employing them to recover the human proteins, obtaining the human proteins and in expressing the proteins via recombinant techniques.

EXAMPLE I

Isolation of Bovine Bone Inductive Factor

Ground bovine bone powder (20-120 mesh, Helitrex) is prepared according to the procedures of M. R. Urist et al., Proc. Natl Acad. Sci USA, 70:3511 (1973) with elimination of some extraction steps as identified below. Ten kgs of the ground powder is demineralized in successive changes of 0.6N HCl at 4°C over a 48 hour period with vigorous stirring. resulting suspension is extracted for 16 hours at 4°C with 50 liters of 2M CaCl2 and 10mM ethylenediamine-tetraacetic acid [EDTA]. and followed by extraction for 4 hours in 50 liters of The residue is washed three times with distilled 0.5M EDTA. water before its resuspension in 20 liters of 4M guanidine hydrochloride [GuCl], 20mM Tris (pH 7.4), 1mM N-ethylmaleimide, 1mM iodoacetamide, 1mM phenylmethylsulfonyl fluorine as described in Clin. Orthop. Rel. Res., 171: 213 (1982). After 16 to 20 hours the supernatant is removed and replaced with another 10 liters of GuCl buffer. The residue is extracted for another 24 hours.

The crude GuCl extracts are combined, concentrated approximately 20 times on a Pellicon apparatus with a 10,000 molecular weight cut-off membrane, and then dialyzed in 50mM Tris, 0.1M NaCl, 6M urea (pH7.2), the starting buffer for the first column. After extensive dialysis the protein is loaded on a 4 liter DEAE cellulose column and the unbound fractions

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are collected.

The unbound fractions are concentrated and dialyzed against 50mM NaAc, 50mM NaCl (pH 4.6) in 6M urea. The unbound fractions are applied to a carboxymethyl cellulose column. Protein not bound to the column is removed by extensive washing with starting buffer, and the bone inductive factor containing material desorbed from the column by 50mM NaAc, 0.25mM NaCl, 6M urea (pH 4.6). The protein from this step elution is concentrated 20- to 40- fold, then diluted 5 times with 80mM KPO₄, 6M urea (pH6.0). The pH of the solution is adjusted to 6.0 with 500mM K₂HPO₄. The sample is applied to an hydroxylapatite column (LKB) equilibrated in 80mM KPO₄, 6M urea (pH6.0) and all unbound protein is removed by washing the column with the same buffer. Bone inductive factor activity is eluted with 100mM KPO₄ (pH7.4) and 6M urea.

The protein is concentrated approximately 10 times, and solid NaCl added to a final concentration of 0.15M. This material is applied to a heparin - Sepharose column equilibrated in 50mM KPO₄, 150mM NaCl, 6M urea (pH7.4). After extensive washing of the column with starting buffer, a protein with bone inductive factor activity is eluted by 50mM KPO₄, 700mM NaCl, 6M urea (pH7.4). This fraction is concentrated to a minimum volume, and 0.4ml aliquots are applied to Superose 6 and Superose 12 columns connected in series, equilibrated with 4M GuCl, 20mM Tris (pH7.2) and the columns developed at a flow rate of 0.25ml/min. The protein demonstrating bone inductive factor activity has a relative migration corresponding to approximately 30,000 dalton protein.

The above fractions are pooled, dialyzed against 50mM NaAc, 6M urea (pH4.6), and applied to a Pharmacia MonoS HR column. The column is developed with a gradient to 1.0M NaCl, 50mM NaAc, 6M urea (pH4.6). Active fractions are pooled and brought to pH3.0 with 10% trifluoroacetic acid (TFA). The material is applied to a 0.46 x 25cm Vydac C4 column in 0.1% TFA and the column developed with a gradient to 90% acetonitrile, 0.1% TFA

(31.5% acetonitrile, 0.1% TFA to 49.5% acetonitrile, 0.1% TFA in 60 minutes at 1ml per minute). Active material is eluted at approximately 40-44% acetonitrile. Aliquots of the appropriate fractions are iodinated by one of the following methods: P. J. McConahey et al, Int. Arch. Allergy, 29:185-189 (1966); A. E. Bolton et al, Biochem J., 133:529 (1973); and D. F. Bowen-Pope, J. Biol. Chem., 237:5161 (1982). The iodinated proteins present in these fractions are analyzed by SDS gel electrophoresis and urea Triton X 100 isoelectric focusing. At this stage, the bone inductive factor is estimated to be approximately 10-50% pure.

EXAMPLE II

Characterization of Bovine Bone Inductive Factor

A. Molecular Weight

Approximately 20ug protein from Example I is lyophilized and redissolved in 1X SDS sample buffer. After 15 minutes of heating at 37°C, the sample is applied to a 15% SDS polyacrylamide gel and then electrophoresed with cooling. molecular weight is determined relative to prestained molecular weight standards (Bethesda Research Labs). Immediately after completion, the gel lane containing bone inductive factor is sliced into 0.3cm pieces. Each piece is mashed and 1.4ml of 0.1% SDS is added. The samples are shaken gently overnight at room temperature to elute the protein. Each gel slice is desalted to prevent interference in the biological assay. The supernatant from each sample is acidified to pH 3.0 with 10% TFA, filtered through a 0.45 micron membrane and loaded on a 0.46cm x 5cm C4 Vydac column developed with a gradient of 0.1% TFA to 0.1% TFA, 90% CH3CN. The appropriate bone inductive factor - containing fractions are pooled and reconstituted with 20mg rat matrix. In this gel system, the majority of bone inductive factor fractions have the mobility of a protein having a molecular weight of approximately 28,000 - 30,000 daltons.

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B. Isoelectric Focusing

The isoelectric point of bone inductive factor activity is determined in a denaturing isoelectric focusing system. Triton X100 urea gel system (Hoeffer Scientific) is modified as follows: 1) 40% of the ampholytes used are Servalyte 3/10; 60% are Servalyte 7-9. 2) The catholyte used is 40mM Approximately 20ug of protein from Example I is lyophilized, dissolved in sample buffer and applied to the isoelectrofocusing gel. The gel is run at 20 watts, 10°C for approximately 3 hours. At completion the lane containing bone inductive factor is sliced into 0.5 cm slices. piece is mashed in 1.0ml 6M urea, 5mM Tris (pH 7.8) and the samples agitated at room temperature. The samples are acidified, filtered, desalted and assayed as described above. The major portion of activity as determined in the assay described in Example III migrates in a manner consistent with a pI of 8.8 - 9.2.

C. Subunit Characterization

The subunit composition of bone inductive factor is also determined. Pure bone inductive factor is isolated from a preparative 15% SDS gel as described above. A portion of the sample is then reduced with 5mM DTT in sample buffer and re-electrophoresed on a 15% SDS gel. The approximately 30kd protein yields two major bands at approximately 20kd and 18kd, as well as a minor band at 30kd. The broadness of the two bands indicates heterogeneity caused most probably by glycosylation, other post translational modification, proteolytic degradation or carbamylation.

EXAMPLE III

Biological Activity of Bone Inductive Factor

A rat bone formation assay according to the general procedure of Sampath and Reddi, Proc. Natl. Acad. Sci. U.S.A.,

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80:6591-6595 (1983) is used to evaluate the osteogenic activity of the bovine bone inductive factor of the present invention obtained in Example I. This assay can also be used to evaluate bone inductive factors of other species. The ethanol precipitation step is replaced by dialyzing the fraction to be assayed against water. The solution or suspension is then redissolved in a volatile solvent, e.g. 0.1 - 0.2 % TFA, and the resulting solution added to 20mg of rat matrix. material is frozen and lyophilized and the resulting powder enclosed in #5 gelatin capsules. The capsules are implanted subcutaneously in the abdominal thoracic area of 21 - 49 day old male long Evans rats. The implants are removed after 7 -14 days. Half of each implant is used for alkaline phosphatase analysis [See, A. H. Reddi et al., Proc. Natl Acad Sci., 69:1601 (1972)] and half is fixed and processed for histological Routinely, lum glycolmethacrylate sections are stained with Von Kossa and acid fuschin to detect new bone Alkaline phosphatase, an enzyme produced by chondroblasts and osteoblasts in the process of matrix formation, is also measured. New cartilage and bone formation often correlates with alkaline phosphatase levels. below illustrates the dose response of the rat matrix samples including a control not treated with bone inductive factor.

TABLE 1

Protein* Implanted ug	<u>Cartilage</u>	Alk. Phos.u/l
7.5	2	Not done
2 5	3	445.7
0.83	3	77.4
0.28	0	32.5
0.00	0	31.0

*At this stage the bone inductive factor is approximately 10-15% pure.

The bone or cartilage formed is physically confined to the space occupied by the matrix. Samples are also analyzed by SDS gel electrophoresis and isoelectric focusing as described above, followed by autoradiography. Analysis reveals a correlation of activity with protein bands at 28 - 30kd and a pI 9.0. An extinction coefficient of 1 OD/mg-cm is used as an estimate for protein and approximating the purity of bone inductive factor in a particular fraction. In the <u>in vivo</u> rat bone formation assays on dilutions as described above, the protein is active <u>in vivo</u> at 10 to 200ng protein/gram bone to probably greater than lug protein/gram bone.

EXAMPLE IV

Bovine Bone Inductive Factor Protein Composition

The protein composition of Example IIA of molecular weight 28 - 30kd is reduced as described in Example IIC and digested with trypsin. Eight tryptic fragments are isolated by standard procedures having the following amino acid sequences:

Fragment 1: A A F L G D I A L D E E D L G

Fragment 2: A F Q V Q Q A A D L

Fragment 3: N Y Q D M V V E G

Fragment 4: S T P A Q D V S R

Fragment 5: N Q E A L R

Fragment 6: L S E P D P S H T L E E

Fragment 7: F D A Y Y

Fragment 8: L K P S N ? A T I Q S I V E

A less highly purified preparation of protein from bovine bone is prepared according to a purification scheme similar to that described in Example I. The purification basically varies from that previously described by omission of the DE-52 column, the CM cellulose column and the mono S column, as well as a reversal in the order of the hydroxylapatite and heparin sepharose columns. Briefly, the concentrated crude 4 M extract is brought to 85% final concentration of ethanol at 4 degrees. The mixture is then centrifuged, and the precipitate redissolved in 50 mM Tris, 0.15 M NaCl, 6.0 M urea. This material is then fractionated on Heparin Sepharose as described. The Heparin bound material

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is fractionated on hydroxyapatite as described. The active fractions are pooled, concentrated, and fractionated on a high resolution gel filtration (TSK 30000 in 6 M guanidinium chloride, 50 mM Tris, pH 7.2). The active fractions are pooled, dialyzed against 0.1% TFA, and then fractionated on a C4 Vydac reverse phase column as described. The preparation is reduced and electrophoresed on an acrylamide gel. The protein corresponding to the 18K band is eluted and digested with trypsin. Tryptic fragments are isolated having the following amino acid sequences:

Fragment 9: S L K P S N H A T I Q S ? V

Fragment 10: S F D A Y Y C S ? A

Fragment 11: V Y P N M T V E S C A

Fragment 12: V D F A D I ? W

Tryptic Fragments 7 and 8 are noted to be substantially the same as Fragments 10 and 9, respectively.

A. bBMP-1

Probes consisting of pools of oligonucleotides (or unique oligonucleotides) are designed according to the method of R. Lathe, <u>J. Mol. Biol.</u>, 183 (1):1-12 (1985) and synthesized on an automated DNA synthesizer. One probe consists of a relatively long (32 nucleotides) "guessmer" [See J. J. Toole et al, <u>Nature</u>, 312:342-347 (1984)] of the following nucleotide sequence:

TCCTCATCCAGGGCAATGTCGCCCAGGAAGGC

Because the genetic code is degenerate (more than one codon can code for the same amino acid), the number of oligonucleotides in a probe pool is reduced based on the frequency of codon usage in eukaryotes, the relative stability of G:T base pairs, and the relative infrequency of the dinucleotide CpG in eukaryotic coding sequences [see Toole et al., supra.]. The second set of probes consists of shorter oligonucleotides (17 nucleotides in length) which contain all possible sequences that could encode the amino acids. The second set of probes has the following sequences:

PCT/US87/01537

- (a) A [A/G] [A/G] TC [T/C] TC [T/C] TC [A/G] TC [T/C] AA
- (b) A [A/G] [A/G] TC [T/C] TC [T/C] TC [A/G] TCNAG

 Bracketed nucleotides are alternatives. "N" means either A,
 T, C or G.

In both cases the regions of the amino acid sequence used for probe design are chosen by avoiding highly degenerate codons where possible. The oligonucleotides are synthesized on an automated DNA synthesizer; the probes are then radioactively labeled with polynucleotide kinase and ³²P-ATP.

These two sets of probes are used to screen a bovine genomic recombinant library. The library is constructed as follows: Bovine liver DNA is partially digested with the restriction endonuclease enzyme Sau 3A and sedimented through a sucrose gradient. Size fractionated DNA in the range of 15-30kb is then ligated to the bacteriophage Bam HI vector EMBL3 [Frischauf et al, J. Mol. Biol., 170:827-842 (1983)]. The library is plated at 8000 recombinants per plate. Duplicate nitrocellulose replicas of the plaques are made and amplified according to a modification of the procedure of Woo et al, Proc. Natl. Acad. Sci. USA, 75:3688-91 (1978).

The 32 mer probe is kinased with 32P-gamma-ATP and hybridized to one set of filters in 5X SSC, 0.1% SDS, 5X Denhardts, 100ug/ml salmon sperm DNA at 45 degrees C and washed with 5X SSC, 0.1% SDS at 45 degrees C. The 17 mer probes are kinased and hybridized to the other set of filters in 3M tetramethylammonium chloride (TMAC), 0.1M sodium phosphate pH6.5, 1mM EDTA, 5X Denhardts, 0.6% SDS, 100ug/ml salmon sperm DNA at 48 degrees C, and washed in 3M TMAC, 50mM Tris pH8.0 at 50 degrees C. These conditions minimize the detection of mismatches to the 17 mer probe pool [see, Wood et al, Proc. Natl. Acad. Sci. U.S.A., 82:1585-1588 (1985)]. recombinants are screened by this procedure and one duplicate positive is plaque purified. DNA is isolated from a plate lysate of this recombinant bacteriophage designated lambda bP-50. bP-50 was deposited December 16, 1986 with the American

Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland USA (hereinafter the "ATCC") under accession number 40295. This deposit as well as the other deposits contained herein meets the requirements of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and Regulations thereunder. This bp-50 clone encodes at least a portion of the bovine bone growth factor designated bBMP-1.

The oligonucleotide hybridizing region of this bBMP-1 clone is localized to an approximately 800bp Eco RI fragment which is subcloned into M13 and sequenced by standard techniques. The partial DNA sequence and derived amino acid sequence of lämbda bP-50 are shown below in Table II. The amino acid sequences corresponding to the tryptic fragments isolated from the bovine bone 28 to 30kd material are underlined in The first underlined portion of the sequence corresponds to tryptic Fragment 1 above from which the oligonucleotide probes are designed. The second underlined portion corresponds to tryptic Fragment 2 above. The predicted amino acid sequence indicates that tryptic Fragment 2 is preceded by a basic residue (R) as expected considering the specificity of trypsin. The nucleic acid sequence preceding the couplet CT at nucleotide positions #292-293 in Table II is presumed to be an intron (noncoding sequence) based on the presence of a consensus acceptor sequence (i.e., a pyrimidine rich tract, TCTCTCTCC, followed by AG) and the lack of a basic residue in the appropriate position of the derived amino acid sequence. This bBMP-1 genomic sequence appears in Table II. The presumptive bBMP-1 peptide sequence from this genomic clone is 37 amino acids in length and is encoded by the DNA sequence from nucleotide #294 through #404 in Table II.

TABLE II

280 290 · (1) 308 323

CCTTGCCTCT TCTCTCCCA GCT GCC TTC CTT GGG GAC ATC GCC CTG GAC GAG GAG

Ala Phe Leu Gly Asp Ile Ala Leu Asp Glu Glu

338 353 368

GAC TTG AGG GCC TTC CAA GTG CAG CAG GCT GCG GAC CTC AGA CAG CGT GCA ACC

Asp Leu Arg Ala Phe Gln Val Gln Gln Ala Ala Asp Leu Arg Gln Arg Ala Thr

383: 398 (37) 414 424 CGC AGG TCT TCC ATC AAA GCT GCA GGTACACTGG GTACAGGCCA Arg Arg Ser Ser Ile Lys Ala Ala

22

B. bBMP-2

Two probes consisting of pools of oligonucleotides are designed on the basis of the amino acid sequence of Fragment 3 and synthesized on an automated DNA synthesizer as described above.

Probe #1: A C N A C C A T [A/G] T C [T/C] T G [A/G] A T
Probe #2: C A [A/G] G A [T/C] A T G G T N G T N G A
These probes are radioactively labeled and employed to screen
the bovine genomic library constructed as described in part A
except that the vector is lambda J1 Bam H1 arms [Mullins et al
Nature 308: 856-858 (1984).] The radioactively labelled 17-mer
Probe #1 is hybridized to the set of filters according to the
method for the 17 mer probe described in part A.

400,000 recombinants are screened by the procedure described above in Part A. One duplicate positive is plaque purified and the DNA is isolated from a plate lysate of the recombinant bacteriophage designated lambda bP-21. Bacteriophage bP-21 was deposited with the ATCC under accession number ATCC 40310 on March 6, 1987. The bP-21 clone encodes the bovine growth factor designated bBMP-2.

The oligonucleotide hybridizing region of this bBMP-2 clone is localized to an approximately 1.2 kb Sac I restriction fragment which is subcloned into M13 and sequenced by standard techniques. The partial DNA sequence and derived amino acid sequence of this Sac I fragment and the contiguous Hind III-Sac I restriction fragment of bP-21 are shown below in Table The bBMP-2 peptide sequence from this clone is 129 amino acids in length and is encoded by the DNA sequence from nucleotide #1 through nucleotide #387. The amino acid sequence corresponding to the tryptic fragment isolated from the bovine bone 28 to 30kd material is underlined in Table III. The underlined portion of the sequence corresponds to tryptic Fragment 3 above from which the oligonucleotide probes for bBMP-2 are designed. The predicted amino acid sequence indicates that tryptic Fragment 3 is preceded by a basic

residue (K) as expected considering the specificity of trypsin. The arginine residue encoded by the CGT triplet is presumed to be the carboxy-terminus of the protein based on the presence of a stop codon (TAG) adjacent to it.

TABLE III

	· · · · · · · · · · · · · · · · · · ·													
(1) GGC G	CAC H	GAT D	GGG G	15 AAA K	GGA G	CAC H		CTC L	30 CAC H	AGA R	AGA R	GAA E	AAG K	45 CGG R
CAA Q	GCA A	AAA K	CAC H	60 AAA K	CAG		AAA K		75 CTC L	AAG K	TCC S	AGC S	TGT C	90 AAG K
AGA R	CAC H		TTA L	105 TAT Y	gtg V	GAC D	TTC F	AGT S	120 GAT D	GTG V	GGG G	TGG W		135 GAC D
TGG W	ATC I	GTT V	GCA A	150 CCG P		GGG G		CAT H				TGC C	CAT H	180 GGG G
GAG E	TGC C	CCT P	TTT F	195 CCC P	CTG L		GAT D	CAC H	210 CTT L		TCC S	ACG T	AAT N	225 CAT H
GCC A	ATT I		CAA Q	240 ACT T	CTG L	GTC V	AAC N	TCA S	255 GTT V	AAC N	TCT S	AAG K	ATT I	270 CCC P
AAG K	GCA A	TGC C	TGT C	385 GTC V	CCA P	ACA T		CTC L	300 AGC S	GCC A	ATC I	TCC S	ATG M	315 CTG L
TAC Y	CTT L	GAT D	GAG E	330 AAT N	GAG E	AAG K	GTG V	GTA V		AAG K	AAC N	TAT Y	CAG Q	360 GAC D
ATG M	GTT V	GTC V	GAG E	375 GGT <u>G</u>		GGG G	TGŤ C		TAGO	CACAC	397 SCA <i>1</i>	\AAT?	40 PAAA)7 !A
417 427 437 447 457 TAAATATATA TATATATATA TTAGAAAAAC AGCAAAAAAA TCAAGTTGAC														
ACTT		67 AT T	TCCC	47 AATG		ACTI	487 TATT		'GGAA	497 ATGG	AATG	5 GAGA	07 AA	
AAGA	_	17 CA C			7 T GA		537 TATA	TTT	'ATAT	547 CTA	CCGA	S AAA	557 SAA	
GTTG		67 AA C	AAAT		7 T AA		587 AGAA		TT			·		

C. bBMP-3

Probes consisting of pools of oligonucleotides are designed on the basis of the amino acid sequences of the tryptic Fragments 9 (Probe #3), 10 (Probe #2), and 11 (Probe #1), and synthesized on an automated DNA synthesizer.

Probe #1: A C N G T C A T [A/G] T T N G G [A/G] T A

Probe #2: C A [A/G] T A [A/G] T A N G C [A/G] T C [A/G] A A

Probe: #3: T G [A/G/T] A T N G T N G C [A/G] T G [A/G] T T

A recombinant bovine genomic library constructed in EMBL3 is screened by the TMAC hybridization procedure detailed above in part A. 400,000 recombinants are screened in duplicate with Probe #1 which has been labeled with ³²P. All recombinants which hybridized to this probe are replated for secondaries. Triplicate nitrocellulose replicas are made of the secondary plates, and amplified as described. The three sets of filters are hybridized to Probes #1, #2 and #3, again under TMAC conditions. One clone, lambda bP-819, hybridizes to all three probes and is plaque purified and DNA is isolated from a plate lysate. Bacteriophage lambda bP-819 was deposited with the ATCC on June 16, 1987 under accession number 40344. This bP-819 clone encodes the bovine bone growth factor designated bBMP-3.

The region of bP-819 which hybridizes to Probe #2 is localized and sequenced. The partial DNA and derived amino acid sequences of this region are shown in Table IVA. The amino acid sequences corresponding to tryptic Fragments 10 and 12 are underlined. The first underlined sequence corresponds to Fragment 12 while the second corresponds to Fragment 10. This region of bP-819, therefore, which hybridizes to Probe #2 encodes at least 111 amino acids. This amino acid sequence is encoded by the DNA sequence from nucleotide #414 through #746.

TABLE IV. A.

383 393 403 413 (1) GAGGAGGAAG CEGTCTACEG GGGTCCTTCT GCCTCTGCAG AAC AAT GAG CTT CCT GGG GCA Asn Asn Glu Leu Pro Gly Ala 443 458 473 GAA TAT CAG TAC AAG GAG GAT GAA GTA TGG GAG GAG AGG AAG CCT TAC AAG ACT Glu Tyr Gln Tyr Lys Glu Asp Glu Val Trp Glu Glu Arg Lys Pro Tyr Lys Thr 518 CTT CAG ACT CAG CCC CCT GAT AAG AGT AAG AAA AAG AAA CAG AGG AAG GGA Leu Gln Thr Gln Pro Pro Asp Lys Ser Lys Asn Lys Lys Lys Gln Arg Lys Gly 578 CCT CAG CAG AAG AGT CAG ACG CTC CAG TIT GAT GAA CAG ACC CTG AAG AAG GCA Pro Gln Gln Lys Ser Gln Thr Leu Gln Phe Asp Glu Gln Thr Leu Lys Lys Ala 623 AGA AGA AAG CAA TGG ATT GAA CCC CGG AAT TGT GCC AGA CGG TAC CTT AAA GTG Arg Arg Lys Gln Trp Ile Glu Pro Arg Asn Cys Ala Arg Arg Tyr Leu Lys <u>Val</u> 668 683 GAC TIC GCA GAT ATT GGC TGG AGC GAA TGG ATT ATT TOC CCC AAG TCC TIC GAT Asp Phe Ala Asp Ile Gly Trp Ser Glu Trp Ile Ile Ser Pro Lys Ser Phe Asp · 713 728 743 (111) GCC TAT TAC TGC TCC GGA GCG TGC CAG TTC CCC ATG CCA AAG GTAGCCATTG Ala Tyr Tyr Cys Ser Gly Ala Cys Gln Phe Pro MET Pro Lys

766 776 786
TITTITGICC TGICCITCCC ATTICCATAG

27

The region of bP-819 which hybridizes to Probe #1 and The partial DNA and derived #3 is localized and sequenced. amino acid sequences of this region are shown in Table IVB. The amino acid sequences corresponding to tryptic Fragments 9 and 11 are underlined. The first underlined sequence corresponds to Fragment 9 while the second underlined sequence corresponds to Fragment 11. The peptide sequence of this region of bP-819 which hybridizes to Probe #1 and #3 is 64 amino acids in length encoded by nucleotide #305 through #493 of The arginine residue encoded by the AGA triplet is presumed to be the carboxy-terminus of the protein based on the presence of a stop codon (TAA) adjacent to it. nucleic acid sequence preceding the couplet TC (positions 305-306) is presumed to be an intron (non-coding sequence) based on the presence of a consensus acceptor sequence (i.e. a pyrimidine-rich stretch, TTCTCCCTTTTCGTTCCT, followed by AG) and the presence of a stop rather than a basic residue in the appropriate position of the derived amino acid sequence.

bBMP-3 is therefore characterized by the DNA and amino acid sequence of Table IV A and Table IV B. The peptide sequence of this clone is 175 amino acids in length and is encoded by the DNA sequence from nucleotide #414 through nucleotide #746 of Table IV A and nucleotide #305 through nucleotide #493 of Table IV B.

TABLE IV. B.

304 (112) 284 294 319 CTAACCIGIG TICICCCITT TOGITCCIAG TCT TIG AAG CCA TCA AAT CAC GCT ACC Ser Leu Lys Pro Ser Asn His Ala Thr 334 349 364 379 ATC CAG AGT ATA GIG AGA GCT GIG GGG GTC GTC CCT GGA ATC CCC GAG CCT TGC <u>Ile Gln Ser Ile Val</u> Arg Ala Val Gly Val Val Pro Gly Ile Pro Glu Pro Cys 409 424 TGT GIG CCA GAA AAG ATG TCC TCA CTC AGC ATC TTA TTC TTT GAT GAA AAC AAG Cys: Val Pro Glu Lys MET Ser Ser Leu Ser Ile Leu Phe Phe Asp Glu Asn Lys 469 AAT GIG GIA CIT AAA GIA TAT CCA AAC AIG ACA GIA GAG TCT TGI GCT TGC AGA Asn Val Val Leu Lys <u>Val Tyr Pro Asn MET Thr Val Glu Ser Cys Ala</u> Cys Arg 513 523

TAACCIGGIG AAGAACICAT CIGGATGCIT AACTCAATCG

29

EXAMPLE V

Human Bone Inductive Factors

A. hBMP-1

Because the bovine and human bone growth factor genes are presumed to be significantly homologous, the bovine bBMP-1 DNA sequence of Table II (or portions thereof) is used as a The 800bp EcoRI probe to screen a human genomic library. fragment of the bovine genomic clone is labeled with $^{32}\mathrm{p}$ by nick-translation. A human genomic library (Toole et al., supra) is plated on 20 plates at 40,000 recombinants per plate. Duplicate nitrocellulose filter replicas are made of each plate and hybridized to the nick-translated probe in 5 X SSC, 5 X Denhardt's, 100ug/ml denatured salmon sperm DNA, 0.1% SDS (the standard hybridization solution) at 50 degrees centigrade for approximately 14 hours. The filters are then washed in 1 X SSC, 0.1% SDS at 50 degrees centigrade and subjected to autoradiography. Five duplicate positives are isolated and plaque purified. DNA is obtained from a plate lysate of one of these recombinant bacteriophage, designated LP-H1. LP-H1 was deposited with the ATCC on March 6, 1987 under accession number 40311. This clone encodes at least a portion of the human genomic bone growth factor called hBMP-1. The hybridizing region of LP-H1 is localized to a 2.5kb XbaI/HindIII restriction fragment.

The partial DNA sequence and derived amino acid sequence of lambda LP-H1 are shown below in Table V. The peptide sequence from this clone is 37 amino acids in length and is encoded by the DNA sequence from nucleotide #3440 through nucleotide #3550. The coding sequence of Table V is flanked by approximately 28 nucleotides (a presumptive 5' noncoding sequence) as well as approximately 19 nucleotides (a presumptive 3' noncoding sequence. A comparison of the bBMP-1 sequence of Table II with the hBMP-1 genomic sequence of Table V indicates the significant homology between the two.

Because the size of coding regions and the positions

of noncoding regions is generally conserved in homologous genes of different species, the locations of the coding and noncoding regions of the bone inductive factor genes may be identified. Regions of homology between the two species' genes, flanked by RNA processing signals at homologous sites, indicate a coding region.

TABLE V

3419 3429 3439 (1) 3454
CAGCCCTGGC TTCTTCTTTT CTCTTTAGCT GCC TTT CTT GGG GAC ATT GCC CTG GAC
Ala Phe Leu Gly Asp Ile Ala Leu Asp

3469 3484 3499 3514

GAA GAG GAC CTG AGG GCC TTC CAG GTA CAG CAG GCT GTG GAT CTC AGA CGG CAC
Glu Glu Asp Leu Arg Ala Phe Gln Val Gln Gln Ala Val Asp Leu Arg Arg His

3529 3544 (37) 3560 3570 ACA GCT CGT AAG TCC TCC ATC AAA GCT GCA GGTAAGCCGG GTGCCAATGG Thr_Ala Arg Lys Ser Ser Ile Lys Ala Ala

A probe specific for the human coding sequence given in Table V is used to identify a human cell line or tissue which synthesizes bone inductive factor. The probe is made according to the following method. Two oligonucleotides having the following sequences:

- (a) GGGAATTCTGCCTTTCTTGGGGACATTGCCCTGGACGAAGAGGACCTGAG
- (b) CGGGATCCGTCTGAGATCCACAGCCTGCTGTACCTGGAAGGCCCTCAGG are synthesized on an automated synthesizer, annealed, extended using the Klenow fragment of E. coli DNA polymerase I, digested with the restriction enzymes Eco RI and Bam HI, and inserted into an M13 vector. A single-stranded 32P-labeled probe is then from template preparation of this subclone by standard techniques. Polyadenylated RNAs from various cell and tissue sources are electrophoresed on formaldehyde-agarose gels and transfered to nitrocellulose by the method of Toole et al., The probe is then hybridized to the nitrocellulose blot in 50% formamide, 5 X SSC, 0.1% SDS, 40 mM sodium phosphate pH 6.5, 100 ug/ml denatured salmon sperm DNA, and 5 mM vanadyl ribonucleosides at 42° C overnight and washed at 65° C in 0.2 X SSC, 0.1% SDS. Following autoradiography, the lane containing RNA from the human osteosarcoma cell line U-2 OS contains hybridizing bands corresponding to RNA species of approximately 4.3 and 3.0 kb.

cDNA is synthesized from U-2 OS polyadenylated RNA and cloned into lambda gt10 by established techniques (Toole et al., <u>supra</u>). 20,000 recombinants from this library are plated on each of 50 plates. Duplicate nitrocellulose replicas are made of the plates. The above described oligonucleotides are kinased with ³²P-gamma-ATP and hybridized to the two sets of replicas at 55° centigrade in standard hybridization solution overnight. The filters are then washed in 1 X SSC, 0.1% SDS at 55° centigrade and subjected to autoradiography. One duplicate positive, designated lambda U2OS-1, is plaque purified. Lambda U2OS-1 was deposited with the ATCC on June 16, 1987 under accession number 40343.

The entire nucleotide sequence and derived amino acid sequence of the insert of lambda U2OS-1 is given in Table VI. This cDNA clone encodes a Met followed by a hydrophobic leader sequence characteristic of a secreted protein, and contains a stop codon at nucleotide positons 2226 - 2228. This clone contains an open reading frame of 2190bp, encoding a protein of 730 amino acids with a molecular weight og 83kd based on this amino acid sequence. The clone contains sequence identical to the coding region given in Table V. This protein is contemplated to represent a primary translation product which is cleaved upon secretion to produce the hBMP-1 protein. This clone is therefore a cDNA for hBMP-1 corresponding to human gene fragment contained in the genomic hBMP-1 sequence lambda LP-H1. noted that amino acids #550 to #590 of BMP-1 are homologous to epidermal growth factor and the "growth factor" domains of Protein C, Factor X and Factor IX.

TABLE VI

															•		
CTA	GAGG	10 555 (CITO		20 3C (X	3003(3(AGC 2					50 SCC (Ala <i>i</i>			
		CIC Leu															
GAC Asp	TAC	ACC Thr	125 TAT Tyr	GAC Asp	CIG Leu	GOG Ala	GAG Glu	140 GAG Glu	GAC Asp	GAC Asp	TCG Ser	GAG Glu	155 CCC Pro	CIC Leu	AAC Asn	TAC Tyr	AAA Lys
170 GAC Asp	rccc Pro	TGC Cys	AAG Lys	GCG Ala	185 GCT Ala	GCC Ala	TTT Phe	CTT Leu	GGG Gly	200 GAC Asp	ATT Ile	GCC Ala	CIG Leu	GAC Asp	215 GAA Glu	GAG Glu	GAC Asp
CIG Leu	AGG Arg	230 GCC Ala	TTC Phe	CAG Gln	GTA Val	CAG Gln	245 CAG Gln	GCT Ala	GIG Val	GAT Asp	CTC Leu	260 AGA Arg	ŒG	CAC His	ACA Thr	GCT Ala	275 OGT Arg
AAG Lys	TCC Ser	TCC Ser	ATC Ile	290 AAA Lys	GCT Ala	GCA Ala	GTT Val	CCA Pro	305 GGA Gly	AAC	ACT Thr	TCT Ser	ACC Thr	320 CCC Pro	AGC Ser	TGC Cys	CAG Gln
AGC Ser	335 ACC Thr	AAC Asn	ccc Gly	CAG Gln	CCT Pro	350 CAG Gln	AGG Arg	GGA Gly	GCC Ala	TGT Cys	365 GGG Gly	AGA Arg	TGG Trp	AGA Arg	GGT Gly	380 AGA Arg	TCC Ser
OGT Arg	AGC Ser	CGG Arg	395 ŒG Ar g	GCG Ala	GCG Ala	ACG Thr	TCC Ser	410 CGA Arg	CCA Pro	GAG Glu	OGT Arg	GIG Val	425 TGG Trp	CCC Pro	GAT Asp	GGG Gly	GIC Val
440 ATC Ile	ccc Pro	TTT Phe	GIC Val	ATT Ile	455 GGG Gly	GGA Gly	AAC Asn	TTC Phe	ACT Thr	470 GGT Gly	AGC Ser	CAG Gln	AGG Arg	GCA Ala	485 GTC Val	TTC Phe	ŒG Arg
CAG Gln	GCC Ala	500 ATG MET	AGG Arg	CAC His	TGG Trp	GAG Glu	515 AAG Lys	CAC His	ACC Thr	TGI Cys	GIC Val	530 ACC Thr	TTC Phe	CIG Leu	CAG Glu	OGC Arg	545 ACT Thr
GAC Asp	GAG Glu	GAC Asp	AGC Ser	560 TAT Tyr	ATT Ile	GIG Val	TTC Phe	ACC Thr	575 TAT Tyr	ŒA Arg	CCT Pro	TGC Cys	GGG Gly	590 TGC Cys	TGC Cys	TCC Ser	TAC Tyr
<i>G</i> IG Val	605 GGT Gly	OGC Arg	œc Arg	GGC Gly	GGG Gly	620 GGC Gly	ccc Pro	CAG Gln	GCC Ala	ATC Ile	635 TCC Ser	ATC Ile	GGC Gly	AAG Lys	AAC Asn	650 TGT Cys	GAC Asp

			665					680					695				
AAG	TTC	GGC	ATT	GIG	GTC	CAC	GAG	CIG	GGC	CAC	GTC	GTC	GGC	TIC	TGG	CAC	GAZ
Lys	Phe	Gly	Ile	Val	Val	His	Glu	Leu	Gly	His	Val	Val	Gly	Phe	Trp	His	Gli
710					725					740			_		755		
CAC	ACT	α	CCA	GAC	CCG	GAC	CCC	CAC	GII	TCC	ATC	GIT	CCI	GAG	AAC	ATC	CAC
His	Thr	Arg	Pro	Asp	Arg	Asp	Arg	His	Val	Ser	Ile	Val	Arg	Glu	Asn	Ile	Gli
		770					785					800					815
			GAG														
Pro	Gly	Gln	Glu	Tyr	Asn	Phe	Leu	Lys	MET	Glu	Pro	GIn	Glu	Val	GIU	ser	Let
				830					845	~~		~~		860	3.03		m~
			TAT														
GLY	GIU	'Inr	Tyr	Asp	Pne	Asp	Ser	TTE	WEI.	HIS	ıyr	ALA	Arg	Asn	unr	me	Sei
	075					000					005					020	
300	875			~	~3.T	890	3000	arra.	~~~	330	905	C3.C	conc.	330	ccc	920	337
			TIC														
Arg	GTĀ	тте	Phe	Teu	ASp	THE	TTG	var	PIO	τλ2	TÄL	GIU	Val	WZII	GTĀ	va.	ту
			935					950					965				
Cuti	\sim	אחחי	GGC		እርር	አ <i>ር</i> አ	CCC.		ACC	ልልር	ccc	GAC		ccc	CAA	ccc	CCC
			Gly														
110	FIO	116	GLY	GHI	my	1111	my		عصد	цуS	OLY	بإحد	116	nu	٠	,,,,,,	
980					995					1010				-	L025	•	
	CIT	TAC	AAG	TGC		GCC	TGT	GGA			CTG	CAA	GAC			GGC	AAC
			Lys														
-		-	•	•			-	•					_			_	
		1040				:	1055					1070				:	1085
TTC	TCC	TCC	CCT	GAA	TAC	α	AAT	GGC	TAC	TCT	GCT	CAC	ATG	CAC	TGC	GIG	TGG
Phe	Ser	Ser	Pro	Glu	Tyr	Pro	Asn	Gly	Tyr	Ser	Ala	His	MET	His	Cys	Val	Tr
			_	L100					1115					130			
			GIC														
Arg	Ile	Ser	Val	Thr	Pro	Gly	Glu	Lys	Ile	Ile	Leu	Asn	Phe	Thr	Ser	Leu	Asp
														•			
	1145					160			~~~		1175	~~	~	~~3	_	1190	mm~
CIG	TAC	CCC.	AGC	CCC	CIG	IGC	TGG	TAC	GAC	TAT.	GIG	GAG:	GIU	U-A	GAT	GGC	TIC
Leu	JĀL	Arg	Ser	Arg	Leu	Cys	пр	ıyr	Asp	JAL	vaı	GIU	vai	Arg	Asp	GTĀ	Me
		,	205				,	220				,	225				
	3.00		.205 GCG	~~	CTICO.	~~3		220	mm~	mcc.	ccc		.235	CTTC	∕~	CNG	COLD.
пр	Arg	TÃ2.	Ala	PIO	Leu	Arg	GTĀ	Arg	Pile	Cys	GTĀ	Ser	туз	Leu	PLO	Giu	FLO
1250	,			1	265				1	280				1	295		
		TYY	ACT ·	_		CCC	CITY.	TCC			ڪليك	~~	ልርረ	-		ייעג	ጥርር
			Thr .														
	7 CAL			- -		9	عانب					9					
	1	310				1	325				7	340				1	.355
بليك			GGC '	יאדני	بلبلت			TAC	GAA	GCC	_	-	GGG	GGT	GAT		
			GIV '														

1385 1400 1370 AAG GAC TAT GGC CAC ATT CAA TOG COC AAC TAC CCA GAC GAT TAC CGG COC AGC Lys Asp Tyr Gly His Ile Gln Ser Pro Asn Tyr Pro Asp Asp Tyr Arg Pro Ser AAA GIC IGC ATC IGG OGG ATC CAG GIG TCI GAG GGC TIC CAC GIG GGC CIC ACA Lys Val Cys Ile Trp Arg Ile Gln Val Ser Glu Gly Phe His Val Gly Leu Thr 1490 1505 TTC CAG TCC TTT CAG ATT CAG CGC CAC GAC AGC TGT GCC TAC CAC TAT CIG CAG Phe Gln Ser Phe Glu Ile Glu Arg His Asp Ser Cys Ala Tyr Asp Tyr Leu Glu 1535 1550 GIG CGC: GAC: GGG CAC AGT GAG AGC AGC ACC CTC ATC GGG CGC TAC TGT GGC TAT Val Arg Asp Gly His Ser Glu Ser Ser Thr Leu Ile Gly Arg Tyr Cys Gly Tyr 1595 1610 GAG AAG CCT GAT GAC ATC AAG AGC AGG TOC AGC CGC CTC TGG CTC AAG TTC GTC Glu Lys Pro Asp Asp Ile Lys Ser Thr Ser Ser Arg Leu Trp Leu Lys Phe Val 1640 1655 1670 TCI GAC GGG TCC ATT AAC AAA GCG GGC TIT GCC GIC AAC TIT TIC AAA GAG GIG Ser Asp Gly Ser Ile Asn Lys Ala Gly Phe Ala Val Asn Phe Phe Lys Glu Val 1685 1700 1715 1730 GAC GAG TGC TCT CGG CCC AAC CGC GGG GGC TGT GAG CAG CGG TGC CTC AAC ACC Asp Glu Cys Ser Arg Pro Asn Arg Gly Cys Glu Gln Arg Cys Leu Asn Thr 1745 1760 1775 CTG GGC AGC TAC AAG TGC AGC TGT GAC CCC GGG TAC GAG CTG GCC CCA GAC AAG Leu Gly Ser Tyr Lys Cys Ser Cys Asp Pro Gly Tyr Glu Leu Ala Pro Asp Lys 1790 1805 1820 OGC OGC TGT GAG GCT GCT TGT GGC GGA TTC CTC ACC AAG CTC AAC GGC TCC ATC Arg Arg Cys Glu Ala Ala Cys Gly Gly Phe Leu Thr Lys Leu Asn Gly Ser Ile 1850: 1865 ACC AGC COG GGC TGG CCC AAG GAG TAC CCC CCC AAC AAG AAC TGC ATC TGG CAG Thr Ser Pro Gly Trp Pro Lys Glu Tyr Pro Pro Asn Lys Asn Cys Ile Trp Gln 1925 CTG GTG GCC CCC ACC CAG TAC CGC ATC TCC CTG CAG TIT GAC TTC TIT GAG ACA Leu Val Ala Pro Thr Gln Tyr Arg Ile Ser Leu Gln Phe Asp Phe Phe Glu Thr 1970 1985 CAG GGC AAT CAT GIG TGC AAG TAC CAC TIC GIG CAG GIG CCC AGI GCA CIC ACA Glu Gly Asn Asp Val Cys Lys Tyr Asp Phe Val Glu Val Arg Ser Gly Leu Thr GCT GAC TOO AAG CTG CAT GGC AAG TTC TGT GGT TCT GAG AAG COO GAG GTC ATC Ala Asp Ser Lys Leu His Gly Lys Phe Cys Gly Ser Glu Lys Pro Glu Val Ile

2060					2075				:	2090					2105			
ACC T	rcc	CAG	TAC	AAC	AAC	ATG	OGC	GIG	GAG	TTC	AAG	TCC	GAC	AAC	ACC	GIG	TCC	
Thr S	er	GIN	TÄL	Asn	ASI	MET.	Arg	vai	GIU	me	гуу	ser	ASD	WZII	TITE	vai	Ser	
	_	120					2135					2150					2165	
AAA A																		
Lys I	Lys	Gly	Phe	Lys	Ala	His	Phe	Phe	Ser	Glu	Lys	Arg	Pro	Ala	Leu	GIN	Pro	
			2	2180				2	2195				:	2210				
CCT. C																		
Pro: A (7	lrg 730)	Gly	Arg	Pro	His	Gln	Leu	Lys	Phe	Arg	Val	Gln	Lys	Arg	Asn	Arg	Thr	
22	225			235		224			2255	-	_	265			275		228	
CCC C		TGA	GCC1	IGC ('AGG(CIC	C GG	:ACCC	CIIC	TIZ	CICA	lgga	ACC.	CACC	TT G	GAC	3GAAT	rg
	22	95		230)5		2315	;	2	2325		23	35		234	5		2355
GGATG	GGG	GC :	PTCGG	TIGOC	C AC	CAAC	20000	CAC	CICC	ACT	CIGO	CATI	.cc (GCCC				GCCC
	23	65		237	75		2385	;	2	395		24	105		241	.5		2425
GACAC	AAC	TG (FIGCI	CICI	T C	10000	ACIG	TGC	CCCFI	2000	CGG2	COGG	GG 1	ACCCI	TCCC	C G	recco	TACC
	24	35		244	15		2455	i	2	2465		24	75		248	15		2495
CCCTC			TGAI						_					GGAC		-	FICCI	

CTAGA

B. hBMP-2: Class I and II

The HindIII-SacI bovine genomic bBMP-2 fragment described in Example IV B. is subcloned into an M13 vector. A ³²P-labeled single-stranded DNA probe is made from a template preparation of this subclone. This probe is used to screen polyadenylated RNAs from various cell and tissue sources as described above in part A. A hybridizing band corresponding to an mRNA species of approximately 3.8 kb is detected in the lane containing RNA from the human cell line U-2 OS. HindIII-SacI fragment is labeled with 32P by nick translation and used to screen the nitrocellulose filter replicas of the above-described U-2 OS cDNA library by hybridization in standard hybridization buffer at 650 overnight followed by washing in 1 X SSC, 0.1% SDS at 650. Twelve duplicate positive clones are picked and replated for secondaries. Duplicate nitrocellulose replicas are made of the secondary plates and both sets hybridized to the bovine genomic probe as the primary screening was performed. One set of filters is then washed in 1 X SSC, 0.1% SDS; the other in 0.1 X SSC, 0.1% SDS at 65°.

Two classes of hBMP-2 cDNA clones are evident based on strong (4 recombinants) or weak (7 recombinants) hybridization signals under the more stringent washing conditions (0.1 X SSC, 0.1% SDS). All 11 recombinant bacteriophage are plaque purified, small scale DNA preparations made from plate lysates of each, and the inserts subcloned into pSP65 and into M13 for sequence analysis. analysis of the strongly hybridizing clones designated hBMP-2 Class I (also known as BMP-2) indicates that they have extensive sequence homology with the sequence given in Table III. These clones are therefore cDNA encoding the human equivalent of the protein encoded by the bBMP-2 gene whose partial sequence is given in Table III. Sequence analysis of the weakly hybridizing recombinants designated hBMP-2 Class II (also known as BMP-4) indicates that they are also quite homologous

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with the sequence given in Table III at the 3' end of their coding regions, but less so in the more 5' regions. Thus they encode a human protein of similar, though not identical, structure to that above.

Full length hBMP-2 Class I cDNA clones are obtained in the following manner. The 1.5 kb insert of one of the Class II subclones (II-10-1) is isolated and radioactively labeled by nick-translation. One set of the nitrocellulose replicas of the U-2 OS cDNA library screened above (50 filters, corresponding to 1,000,000 recombinant bacteriophage) are rehybridized with this probe under stringent conditions (hybridization at 65° in standard hybridization buffer; washing at 650 in 0.2 X SSC, 0.1% SDS). All recombinants which hybridize to the bovine genomic probe which do not hybridize to the Class II probe are picked and plaque purified (10 recombinants). Plate stocks are made and small scale bacteriophage DNA preparations made. After subcloning into M13, sequence analysis indicates that 4 of these represent clones which overlap the original Class I clone. One of these, lambda U2OS-39, contains an approximately 1.5 kb insert and was deposited with the ATCC on June 16, 1987 under accession number 40345. The partial DNA sequence (compiled from lambda U2OS-39 and several other hBMP-2 Class I cDNA recombinants) and derived amino acid sequence are shown below in Table VII. Lambda U20S-39 is expected to contain all of the nucleotide sequence necessary to encode the entire human counterpart of the protein BMP-2 Class II encoded by the bovine gene segment whose partial sequence is presented in Table III. This human cDNA hBMP-2 Class II contains an open reading frame of 1188 bp, encoding a protein of 396 amino acids. This protein of 396 amino acids has a molecular weight of 45kd based on this amino acid sequence. It is contemplated that this sequence represents the primary translation product. The protein is preceded by a 5' untranslated region of 342 bp with stop codons in all frames.

The 13 bp region preceding this 5' untranslated region represents a linker used in the cDNA cloning procedure.

TABLE VII

GICGACICIE		20 GIGIGI	CAGCA	30 CTTG		GGGG	40 ACIT		GAAC	50 TTG	CAGG		60 AT A	ACTIO	70 GOGCA
80 CCCCACTITIO		90 GGIGCC	TTTGC	100 CCCA(GAGO	110 CIGC			120 ICT	COGA		30 CA C	acca	140 CCICC
150 ACTOCTOGGO		160 000GAC	ACTGA		O C TG		180 CAGC			190 GAG	AGAC		00 OG G	ccccc	210 ACCC
220 GGGAGAAGGZ		230 GCAAAG	AAAAG	240 GAAC		CATT	250 CGT			260 CCA	GGTO		70 GA C	CAGAG	280 77777
290 TCCATGIGGA		300 CITICA	ATGGA	310 ŒIGI		ccca	320 FIGC			330 ACG (GACT	_	40 IC T	CCTAA	350 AGGT
(1) CGACC ATG MET							ea go					∞ α			
		_													
CTC CTG GG		GOG GO	r GGC	CTC	430 GIT	ccc	GAG	CIG	GGC	445 CGC	AGG				
Leu Leu Gl	C GGC	GOG GO Ala Al	r GGC a Gly	CTC	430 GIT	ccc	GAG Glu	CIG	GGC	445 CGC	AGG Arg	Lys			
	C GGC Y Gly	GOG GO Ala Al 47. GGC OG	r ccc a Gly 5 c ccc	CTC Leu TCA	430 GIT Val	CCG Pro CAG	GAG Glu 490 CCC	CTG Leu TCT	GGC Gly GAC	445 CGC Arg	AGG Arg GIC	Lys 505 CTG	Phe AGC	Ala	
Leu Leu Gl 460 GCG GCG TC Ala Ala Se	c GGC y Gly c TOG r Ser	GCG GC Ala Al 47 GGC CG Gly An	r GGC a Gly 5 c ccc g Pro	CTC Leu TCA Ser	430 GIT Val TCC Ser	CCG Pro CAG Gln	GAG Glu 490 CCC Pro	CIG Leu TCT Ser	GGC Gly GAC Asp 550	445 CGC Arg CAG Glu	AGG Arg GIC Val	Lys 505 CTG Leu	Phe AGC Ser	GAG Glu 565	
Leu Leu Gl 460 GCG GCG TC Ala Ala Se	oc GGC TOG TOG TSer O G OGG	GCG GC Ala Al 47. GGC CG Gly An	F GGC a Gly	CTC Leu TCA Ser 535 ATG	430 GTT Val TCC Ser	CCG Pro CAG Gln	GAG Glu 490 CCC Pro	CIG Leu TCI Ser	GGC Gly GAC Asp 550 CAG	445 CGC Arg GAG Glu AGA	AGG Arg GIC Val	Lys 505 CTG Leu ACC	Phe AGC Ser	GAG Glu 565 AGC	
Leu Leu Gl 460 GCG GCG TC Ala Ala Se TTC GAG TT	C GGC Y GLY G TOG T Ser O G CGG U Arg	GCG GCC Ala Al 47. GGC CGGLY An CTG CTG Lett Lett 580 GTG CCG	F GGC AGC Pro	CTC Leu TCA Ser 535 ATG MET	430 GIT Val TCC Ser TTC Phe	CCG Pro CAG Gln GGC Gly 595 CTA	GAG Glu 490 CCC Pro CIG Leu	CTG Leu TCT Ser AAA Lys	GGC Gly GAC Asp 550 CAG Gln	445 CGC Arg CAG Glu AGA Arg	AGG Arg GIC Val CCC Pro 610 AGG	Lys 505 CTG Leu ACC Thr	Phe AGC Ser CCC Pro	Ala GAG Glu 565 AGC Ser	
Leu Leu Gl 460 GCG GCG TC Ala Ala Se TTC GAG TT Phe Glu Le AGG GAC GC	C GGC Y GLY TO TOS TO SET O CGG U Arg C GIG TO TOA	GCG GCC Ala Al 47. GGC CGG Gly An CTG CTG Leu Leu 580 GTG CCC Val Pro	F GGC A GIY C CCC Pro C AGC Ser C CCC Pro 640 C CCA	CTC Leu TCA Ser 535 ATG MET TAC Tyr	430 GIT Val TCC Ser TTC Phe ATG MET	CAG Gln GGC Gly 595 CTA Leu	GAG Glu 490 CCC Pro CTG Leu GAC Asp	CIG Leu TCT Ser AAA Lys CIG Leu 655 GAG	GGC Gly GAC Asp 550 CAG Gln TAT Tyr	445 CGC Arg CAG Glu AGA Arg CGC Arg	AGG Arg GIC Val CCC Pro 610 AGG Arg	Lys 505 CTG Leu ACC Thr CAC His	Phe AGC Ser CCC Pro TCG Ser 670 CGA	GAG Glu 565 AGC Ser GGT Gly	

730 745 760 AGT GGG AAA ACA ACC CGG AGA TIC TIT TIT AAT TITA AGT TCT ATC CCC ACG GAG Ser Gly Lys Thr Thr Arg Arg Phe Phe Asn Leu Ser Ser Ile Pro Thr Glu 805 820 835 CAG TIT ATC ACC TCA GCA GAG CIT CAG GII TIC CGA GAA CAG ATG CAA GAT GCI Glu Phe Ile Thr Ser Ala Glu Leu Gln Val Phe Arg Glu Gln MET Gln Asp Ala 850 865 880 TTA GGA AAC AAT AGC AGT TTC CAT CAC CGA ATT AAT ATT TAT GAA ATC ATA AAA Leu Gly Asn Asn Ser Ser Phe His His Arg Ile Asn Ile Tyr Glu Ile Ile Lys 895 910 CCT GCA ACA GCC AAC TOG AAA TTC CCC GTG ACC AGT CIT TTG GAC ACC AGG TTG Pro Ala Thr Ala Asn Ser Lys Phe Pro Val Thr Ser Leu Leu Asp Thr Arg Leu 970 GIG AAT CAG AAT GCA AGC AGG TGG GAA AGT TIT GAT GIC ACC CCC GCI GIG ATG Val Asn Gln Asn Ala Ser Arg Trp Glu Ser Phe Asp Val Thr Pro Ala Val MET 1000 1015 1030 CGG TGG ACT GCA CAG GGA CAC GCC AAC CAT GGA TTC GTG GTG GAA GTG GCC CAC Arg Trp Thr Ala Gln Gly His Ala Asn His Gly Phe Val Val Glu Val Ala His 1075 TIG GAG GAG AAA CAA GGI GIC TOO AAG AGA CAT GIT AGG ATA AGC AGG TOT TIG Leu Glu Glu Lys Gln Gly Val Ser Lys Arg His Val Arg Ile Ser Arg Ser Leu CAC CAA GAT GAA CAC AGC TGG TCA CAG ATA AGG CCA TTG CTA GTA ACT TTT GGC His Gln Asp Glu His Ser Trp Ser Gln Ile Arg Pro Leu Leu Val Thr Phe Gly 1180 1195 CAT GAT GGA AAA GGG CAT CCT CTC CAC AAA AGA GAA AAA CGT CAA GCC AAA CAC His Asp Gly Lys Gly His Pro Leu His Lys Arg Glu Lys Arg Gln Ala Lys His 1240 AAA CAG CCG AAA CCC CIT AAG TCC AGC TGT AAG AGA CAC CCT TTG TAC GTG GAC Lys Gln Arg Lys Arg Leu Lys Ser Ser Cys Lys Arg His Pro Leu Tyr Val Asp 1285 1300 -1315 TTC AGT GAC GIG GGG TGG AAT GAC TGG ATT GIG GCT CCC CCG GGG TAT CAC GCC Phe Ser Asp Val Gly Trp Asn Asp Trp Ile Val Ala Pro Pro Gly Tyr His Ala 1330 1345 1360 1375 TIT TAC TGC CAC GGA GAA TGC CCT TIT CCT CTG GCT GAT CAT CTG AAC TCC ACT Phe Tyr Cys His Gly Glu Cys Pro Phe Pro Leu Ala Asp His Leu Asn Ser Thr 1390 1405 1420 AAT CAT GOC ATT GIT CAG ACG TIG GIC AAC TOT GIT AAC TOT AAG AIT OOT AAG Asn His Ala Ile Val Gln Thr Leu Val Asn Ser Val Asn Ser Lys Ile Pro Lys

1435 1450 1465 1480 GCA TGC TGT GTC CCG ACA GAA CTC AGT GCT ATC TCG ATG CTG TAC CTT GAC GAG Ala Cys Cys Val Pro Thr Glu Leu Ser Ala Ile Ser MET Leu Tyr Leu Asp Glu

1495 1510 1525

AAT GAA AAG GIT GTA TTA AAG AAC TAT CAG GAC ATG GIT GIG GAG GGI TGT GGG
Asn Glu Lys Val Val Leu Lys Asn Tyr Gln Asp MET Val Val Glu Gly Cys Gly

1540(396) 1553 1563 1573 1583 1593 1603 TGT CGC TAGTACAGCA AAATTAAATA CATAAATATA TATATATATA TATATTTTAG AAAAAAGAAA Cys÷Arg:

AAAA .

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Full-length hBMP-2Class II human cDNA clones are obtained in the following manner. The 200 bp EcoRI-SacI fragment from the 5' end of the Class II recombinant II-10-1 is isolated from its plasmid subclone, labeled by nick-translation, and hybridized to a set of duplicate nitrocellulose replicas of the U-2 OS cDNA library (25 filters/set; representing 500,000 recombinants). Hybridization and washing are performed under stringent conditions as described above. 16 duplicate positives are picked and replated for secondaries. Nitrocellulose filter replicas of the secondary plates are made and hybridized to an oligonucleotide which was: synthesized to correspond to the sequence of II-10-1 and is of the following sequence:

CGGGCGCTCAGGATACTCAAGACCAGTGCTG

Hybridization is in standard hybridization buffer AT 50° C with washing at 50° in 1 X SSC, 0.1% SDS. 14 recombinant bacteriophage which hybridize to this oligonucleotide are plaque purified. Plate stocks are made and small scale bacteriophage DNA preparations made. After sucloning 3 of these into M13, sequence analysis indicates that they represent clones which overlap the original Class II clone. these, lambda U2OS-3, was deposited with the ATCC under accession number 40342 on June 16, 1987. U20S-3 contains an insert of approximately 1.8 kb. The partial DNA sequence and derived amino acid sequence of U20S-3 are shown below in This clone is expected to contain all of the Table VIII. nucleotide sequence necessary to encode the entire human BMP-2 Class II protein. This cDNA contains an open reading frame of 1224 bp, encoding a protein of 408 amino acids, preceded by a.5' untranslated region of 394 bp with stop codons in all frames, and contains a 3' untranslated region of 308 bp following the in-frame stop codon. The 8 bp region preceding the 5' untranslated region represents a linker used in the cDNA cloning procedure. This protein of 408 amino acids has molecular weight of 47kd and is contemplated to represent the

primary translation product.

TABLE VIII

10 20 30 40 50 60 CTCTAGAGGG CAGAGGAGGA GGGAGGGAGG GAAGGAGGGC GGAGGCGGGC CGGAAGCTA GGTGAGTG	70 FIG
80 90 100 110 120 130 1 GCATCOGAGC TGAGGGAGGC GAGCCTGAGA CGCCGGCTGCT GCTCCGGCTG AGTATCTAGC TTGTCTCC	.40 CC
150 160 170 180 190 200 2 GATGGGATTC COGTCCAAGC TATCTCGAGC CTGCAGCGCC ACAGTCCCCC GCCCTCGCCC AGGTTCAC	10 TG
220 230 240 250 260 270 2 CAACOGITCA GAGGICCCA GGAGCIGCIG CIGGGGAGCC CGCIACIGCA GGGACCIATG GAGCCATI	80
290 300 310 320 330 340 3 GTAGTGCCAT CCCGAGCAAC GCACTGCTGC AGCTTCCCTG AGCCTTTCCA GCAAGTTTGT TCAAGATT	50 GG
360 370 380 390 400 (1) CIGICAAGAA TCATGGACIG TTATTATATG CCTTGTTTTC TGTCAAGACA CC ATG ATT CCT MET Ile Pro	-
417 432 447 462 GGT AAC CGA ATG CTG ATG GTC GTT TTA TTA TGC CAA GTC CTG CTA GGA GGC GCG Gly Asn Arg MET Leu MET Val Val Leu Leu Cys Gln Val Leu Leu Gly Gly Ala	
477 492 507 AGC CAT GCT AGT TTG ATA CCT GAG ACG GGG AAG AAA AAA GTC GCC GAG ATT CAG Ser His Ala Ser Leu Ile Pro Glu Thr Gly Lys Lys Lys Val Ala Glu Ile Gln	
522 537 552 567 GGC CAC GGG GGA GGA CGC CGC TCA GGG CAG AGC CAT GAG CTC CTG CGG GAC TTC Gly His Ala Gly Gly Arg Arg Ser Gly Gln Ser His Glu Leu Leu Arg Asp Phe	
582 597 612 627 GAG GCG ACA CTT CTG CAG ATG TTT GGG CTG CGC CGC CGC CAG CCT AGC AAG Glu Ala Thr Leu Leu Gln MET Phe Gly Leu Arg Arg Pro Gln Pro Ser Lys	
642 657 672 AGT GCC GTC ATT CCG GAC TAC ATG CGG GAT CTT TAC CGG CTT CAG TCT GGG GAG Ser Ala Val Ile Pro Asp Tyr MET Arg Asp Leu Tyr Arg Leu Gln Ser Gly Glu	
687 702 717 732 GAG GAG GAA GAG CAG ATC CAC AGC ACT CGT CIT GAG TAT CCT GAG CGC CCG GCC Glu Glu Glu Glu Gln Ile His Ser Thr Gly Leu Glu Tyr Pro Glu Arg Pro Ala	

				: ACC	GIG Val				CAC					CIG			
	A GGC				807 A AAC 1 Asn	TCI					CIC					AGC	
			GAG		ATC			GCA					TIC				
					TGG Trp					CAC					TAT		
ATG MEI	957 AAG Lys	000	CCA Pro	GCA Ala	GAA Glu	972 GIG Val	GIG Val	CCT Pro	GGG Gly	CAC His	987 CTC Leu	ATC	ACA Thr	OGA Arg	CTA	1002 CIG Leu	GAC
ACG Thr	AGA Arg	CIG	1017 GIC Val	CAC	CAC His	AAT Asn	GIG	1032 ACA Thr	OGG Arg	TGG Trp	GAA Glu	ACT	1047 TTT Phe	GAT Asp	GIG Val	AGC Ser	CCT Pro
	GIC			TGG	1077 ACC Thr				CAG					CTA			
GIG	ACT	1122 CAC	crc	CAT	CAG Gln] ACT	1137 CGG	ACC	CAC	CAG	GGC	L152 CAG	CAT	GIC	AGG	I ATT	L167 AGC
CGA	TŒ	TTA	CCT	L182 CAA	GGG	AGT	GGG] AAT	.197 TGG	GCC	CAG	CIC] CGG	.212 CCC	CIC	CIG	GIC
:	1227					242]	L257		•	•	נ	272	
Thr	Phe	Gly	His L287	Asp	Gly	Arg	Gly 1	His .302	Ala	Leu	Thr	Arg 1	Arg .317	Arg	Arg	Ala	Lys
	Ser			His	CAC His				Ala					Lys			
α C	CAC			TAT	GIG (GAT	GIG				GAC	TGG		
~~-		392					407					422					437
					CAG (Gln)												

GCT GAC CAC CTC AAC TCA ACC AAC CAT GCC ATT GTG CAG ACC CTG GTC AAT TCT Ala Asp His Leu Asn Ser Thr Asn His Ala Ile Val Gln Thr Leu Val Asn Ser GIC AAT TOO AGT ATO COO AAA GOO TGT TGT GTG COO ACT GAA CTG AGT GOO ATO Val Asn Ser Ser Ile Pro Lys Ala Cys Cys Val Pro Thr Glu Leu Ser-Ala Ile TCC ATG CTG TAC CTG GAT GAG TAT GAT AAG GTG GTA CTG AAA AAT TAT CAG GAG Ser MET Leu Tyr Leu Asp Glu Tyr Asp Lys Val Val Leu Lys Asn Tyr Gln Glu (408)ATG GTA GTA GAG GGA TGT GGG TGC CGC TGAGATCAGG CAGTCCTTGA GGATAGACAG MET Val Val Glu Gly Cys Gly Cys Arg ACAGACIGCT TOCTTATAGC TGGACTTTTA TTTAAAAAAA AAAAAAAAA AATGGAAAAA ATCCCTAAAC ATTCACCITG ACCITATITA TGACITTACG TGCAAATGIT TTGACCATAT TGATCATATA TTTTGACAAA ATATATTTAT AACTACGTAT TAAAAGAAAA AAATAAAATG AGTCATTATT TTAAAAAAAA AAAAAAAACT

CTAGAGIOGA OGGAATIC

The sequences of BMP-2 Class I and II, as well as BMP-3 as shown in Tables III, IV, VII and VIII have significant homology to the beta (B) and beta (A) subunits of the inhibins. The inhibins are a family of hormones which are presently being investigated for use in contraception. See, A. J. Mason et al, Nature, 318:659-663 (1985). To a lesser extent they are also homologous to Mullerian inhibiting substance (MIS), a testicular glycoprotein that causes regression of the Mullerian duct during development of the male embryo and transforming growth factor-beta (TGF-b) which can inhibit or stimulate growth of cells or cause them to differentiate. Furthermore, the sequence of Table VII encoding hBMP-2 Class II has significant homology to the <u>Drosophila</u> decapentaplegic (DPP-C) locus transcript. See, J. Massague, Cell, 49:437-438 (1987); R. W. Padgett et al, Nature, 325:81-84 (1987); R. L. Cate et al, Cell 45: 685-698 (1986). It is considered possible therefore that BMP-2 Class II is the human homolog of the protein made from this transcript from this developmental mutant locus.

C. <u>BMP-3</u>

Because bovine and human bone growth factor genes are presumed to be significantly homologous, oligonucleotide probes which have been shown to hybridize to the bovine DNA sequence of Table IV.A and IV.B are used to screen a human genomic library A human genomic library (Toole et al., supra) is screened using these probes, and presumptive positives are isolated and DNA sequence obtained as described above. Evidence that this recombinant encodes a portion of the human bone inductive factor molecule relies on the bovine/human protein and gene structure homologies.

Once a recombinant bacteriophage containing DNA encoding a portion of the human BMP-3 molecule is obtained the human coding sequence is used as a probe as described in Example V (A) to identify a human cell line or tissue which synthesizes BMP-3. mRNA is selected by oligo (dT) cellulose

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chromatography and cDNA is synthesized and cloned in lambda gt10 by established techniques (Toole et al., supra).

Alternatively, the entire gene encoding this human bone inductive factor can be identified and obtained in additional recombinant clones if necessary. recombinants containing further 3' or 5' regions of this human bone inductive factor gene can be obtained by identifying unique DNA sequences at the end(s) of the original clone and using these as probes to rescreen the human genomic library. The gene can then be reassembled in a single plasmid by standard molecular biology techniques and amplified in The entire human BMP-3 factor gene can then be bacteria. transferred to an appropriate expression vector. The expression vector containing the gene is then transfected into a mammalian cell, e.g. monkey COS cells, where the human gene is transcribed and the RNA correctly spliced. Media from the transfected cells are assayed for bone inductive factor activity as described herein as an indication that the gene is complete. mRNA is obtained from these cells and cDNA synthesized from this mRNA source and cloned. The procedures described above may similarly be employed to isolate other species' bone inductive factor of interest by utilizing the bovine bone inductive factor and/or human bone inductive factor as a probe source. Such other species' bone inductive factor may find similar utility in, inter alia, fracture repair.

EXAMPLE VI

Expression of Bone Inductive Factors.

In order to produce bovine, human or other mammalian bone inductive factors, the DNA encoding it is transferred into an appropriate expression vector and introduced into mammalian cells by conventional genetic engineering techniques.

One skilled in the art can construct mammalian expression vectors by employing the sequence of Tables II-

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VIII or other modified sequences and known vectors, such as pCD [Okayama et al., Mol. Cell Biol., 2:161-170 (1982)] and pJL3; pJL4 [Gough et al., EMBO J., 4:645-653 (1985)]. transformation of these vectors into appropriate host cells can result in expression of osteoinductive factors. skilled in the art could manipulate the sequences of Tables II-VIII by eliminating or replacing the mammalian regulatory sequences flanking the coding sequence with bacterial sequences to create bacterial vectors for intracellular or extracellular expression by bacterial cells. For example, the coding sequences could be further manipulated (e.g. ligated to other known linkers or modified by deleting non-coding sequences there-from or altering nucleotides therein by other known techniques). The modified bone inductive factor coding sequence could then be inserted into a known bacterial vector using procedures such as described in T. Taniguchi et al., Proc. Natl Acad. Sci. USA, 77:5230-5233 (1980). This exemplary bacterial vector could then be transformed into bacterial host cells and bone inductive factor expressed thereby. a strategy for producing extracellular expression of bone inductive factor in bacterial cells., see, e.g. European patent application EPA 177,343.

Similar manipulations can be performed for the construction of an insect vector [See, e.g. procedures described in published European patent application 155,476] for expression in insect cells. A yeast vector could also be constructed employing yeast regulatory sequences for intracellular or extracellular expression of the factors of the present invention by yeast cells. [See, e.g., procedures described in published PCT application WO86/00639 and European patent application EPA 123,289].

A method for producing high levels of an osteoinductive factor of the invention from mammalian cells involves the construction of cells containing multiple copies of the heterologous bone inductive factor gene. The heterologous gene can be linked to an amplifiable marker, e.g. the dihydrofolate reductase (DHFR) gene for which cells containing increased gene copies can be selected for propagation in increasing concentrations of methotrexate (MTX) according to the procedures of Kaufman and Sharp, <u>J. Mol. Biol.</u>, 159:601-629 (1982). This approach can be employed with a number of different cell types.

For example, a plasmid containing a DNA sequence for a bone inductive factor of the invention in operative association with other plasmid sequences enabling expression thereof and the DHFR expression plasmid pAdA26SV(A)3 [Kaufman and Sharp, Mol. Cell. Biol., 2:1304 (1982)] can be co-introduced into DHFR-deficient CHO cells, DUKX-BII, by calcium phosphate coprecipitation and transfection. DHFR expressing transformants are selected for growth in alpha media with dialyzed fetal calf serum, and subsequently selected for amplification by growth in increasing concentrations of MTX (sequential steps in 0.02, 0.2, 1.0 and 5uM MTX) as described in Kaufman et al., Mol Cell Biol., 5:1750 (1983). Transformants are cloned, and biologically active bone inductive factor expression is monitored by rat bone formation assay. Bone inductive factor expression should increase with increasing levels of MTX Similar procedures can be followed to produce resistance. other bone inductive factors.

Alternatively, the human gene is expressed directly, as described above. Active bone inductive factor may be produced in bacteria or yeast cells. However the presently preferred expression system for biologically active recombinant human bone inductive factor is stably transformed CHO cells.

As one specific example, to produce the human bone inductive factor (hBMP-1) of Example V, the insert of U2OS-1 is released from the vector arms by digestion with Sal I and subcloned into the mammalian expression vector pMT2CX digested with Xho I. Plasmid DNA from this subclone is transfected into COS cells by the DEAE-dextran procedure [Sompayrac and

Danna <u>PNAS</u> 78:7575-7578 (1981); Luthman and Magnusson, <u>Nucl.Acids Res</u>. 11: 1295-1308 (1983)]. Serum-free 24 hr. conditioned medium is collected from the cells starting 40-70 hr. post-transfection.

The mammalian expression vector pMT2 Cla-Xho (pMT2 CX) is a derivative of p91023 (b) (Wong et al., Science 228:810-815, 1985) differing from the latter in that it contains the ampicillin resistance gene in place of the tetracycline resistance gene and further contains a XhoI site for insertion of cDNA clones. The functional elements of pMT2 Cla-Xho have been described (Kaufman, R.J., 1985, Proc. Natl. Acad. Sci. USA 82:689-693) and include the adenovirus VA genes, the SV40 origin of replication including the 72 bp enhancer, the adenovirus major late promoter including a 5' splice site and the majority of the adenovirus tripartite leader sequence present on adenovirus late mRNAs, a 3' splice acceptor site, a DHFR insert, the SV40 early polyadenylation site (SV40), and pBR322 sequences needed for propagation in E. coli.

Plasmid pMT2 Cla-Xho is obtained by EcoRI digestion of pMT2-VWF, which has been deposited with the American Type Culture Collection (ATCC), Rockville, MD (USA) under accession number ATCC 67122. EcoRI digestion excises the cDNA insert present in pMT2-VWF, yielding pMT2 in linear form which can be ligated and used to transform <u>E. coli</u> HB 101 or DH-5 to ampicillin resistance. Plasmid pMT2 DNA can be prepared by conventional methods. pMT2CX is then constructed by digesting pMT2 with Eco RV and XbaI, treating the digested DNA with Klenow fragment of DNA polymerase I, and ligating Cla linkers (NEBiolabs, CATCGATG). This removes bases 2266 to 2421 starting from the Hind III site near the SV40 origin of replication and enhancer sequences of pMT2. Plasmid DNA is then digested with EcoRI, blunted as above, and ligated to an EcoRI adapter,

^{5&#}x27; POA-AATTCCTCGAGAGCT 3'

3' GGAGCTCTCGA 5'

digested with XhoI, and ligated, yielding pMT2 Cla-Xho, which may then be used to transform <u>E</u>. <u>coli</u> to ampicillin resistance. Plasmid pMT2 Cla-Xho DNA may be prepared by conventional methods.

Example VII

Biological Activity of Expressed Bone Inductive Factor A. BMP-1

To measure the biological activity of the expressed bone inductive factor (hBMP-1) obtained in Example VI above. The factor is partially purified on a Heparin Sepharose column. 4 ml of transfection supernatant from one 100 mm dish is concentrated approximately 10 fold by ultrafiltration on a YM 10 membrane and then dialyzed against 20mM Tris, 0.15 M NaCl, pH 7.4 (starting buffer). This material is then applied to a 1.1 ml Heparin Sepharose column in starting buffer. Unbound proteins are removed by an 8 ml wash of starting buffer, and bound proteins, including BMP-1, are desorbed by a 3-4 ml wash of 20 mM Tris, 2.0 M NaCl, pH 7.4.

The proteins bound by the Heparin column are concentrated approximately 10-fold on a Centricon 10 and the salt reduced by diafiltration with 0.1% trifluoroacetic acid. The appropriate amount of this solution is mixed with 20 mg of rat matrix and then assayed for in vivo bone and cartilage formation as previously described in Example III. A mock transfection supernatant fractionation is used as a control.

The implants containing rat matrix to which specific amounts of human BMP-1 have been added are removed from rats after seven days and processed for histological evaluation. Representative sections from each implant are stained for the presence of new bone mineral with von Kossa and acid fuschin, and for the presence of cartilage-specific matrix formation using toluidine blue. The types of cells present within the section, as well as the extent to which these cells display phenotype are evaluated.

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Addition of human BMP-1 to the matrix material resulted in formation of cartilage-like nodules at 7 days post implantation. The chondroblast-type cells were recognizable by shape and expression of metachromatic matrix. The amount of activity observed for human BMP-1 was dependent upon the amount of human BMP-1 protein added to the matrix. Table IX illustrates the dose-response relationship of human BMP-1 protein to the amount of bone induction observed.

Table IX

IMPLANT NUMBER	AMOUNT USED (equivalent of mitransfection media)	HISTOLOGICAL SCORE
876-134-1	10 BMP-1	C+2
876-134-2	3 BMP-1	C+1
876-134-3	1 BMP-1	C +/-
876-134-4	10 MOCK	c -
876-134-5	з моск	c -
876-134-6	1 MOCK	c -

Cartilage (c) activity was scored on a scale from O(-) to 5.

Similar levels of activity are seen in the Heparin Sepharose fractionated COS cell extracts. Partial purification is accomplished in a similar manner as described above except that 6 M urea is included in all the buffers. Further, in a rat bone formation assay as described above, BMP-2 has similarly demonstrated chondrogenic activity.

The procedures described above may be employed to isolate other bone inductive factors of interest by utilizing the bovine bone inductive factors and/or human bone inductive factors as a probe source. Such other bone inductive factors may find similar utility in, inter alia, fracture repair.

The foregoing descriptions detail presently preferred embodiments of the present invention. Numerous modifications

and variations in practice thereof are expected to occur to those skilled in the art upon consideration of these descriptions. Those modifications and variations are believed to be encompassed within the claims appended hereto.

Form PCT.RO 134 (January 1981)

		International Application	No: PCT/ /
	MIC	CROORGANISMS	
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A. IDENTIFICATION OF DEI	OSIT		
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Name of depositary institution 4			·
	A continue To		
	American ly	pe Culture Collection	
Address of depositary institution	(including postal code	and country) 4	
	12301 Parkl	awn Drive	
	Rockville,	Maryland 20852 USA	
Name of De <u>posit</u>	ATCC No.	Referred to on	Date of Deposit
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LP-H1	40311	29/20	March 4, 1987
ъР50	40295	20/3	December 15, 198
bP-21	40310	22/18	March 4, 1987
U20S-3	40342	44/22	June 16, 1987
Lambda U2-OS-	-1 40343	32/33	June 16, 1987
Lambda BP819	40344	25/23	June 16, 1987
U20S-39	40345	39/21	June 16, 1987
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WHAT IS CLAIMED IS:

- 1.. A pharmaceutical composition comprising a protein selected from the group consisting of:
 - (a) BMP-1;
 - (b) BMP-2 Class I;
 - (c) BMP-2 Class II;
 - (d) BMP-3; and

mixtures: thereof, in a pharmaceutically acceptable vehicle.

- 22. A composition of Claim 1 wherein said protein is BMP1..
- 3. A composition of Claim I wherein said protein is BMP-2. Class I.
- 4. A composition of Claim 1 wherein said protein is BMP-2 Class II.
- 5. A composition of Claim 1 wherein said protein is BMP3.
- 6. The pharmaceutical composition of Claim 1 further comprising a matrix capable of delivering the composition to the site of the bone defect and providing a structure for inducing bone formation.
- 7. The composition of Claim 6 wherein said matrix comprises a material selected from the group consisting of hydroxyapatite, collagen, polylactic acid and tricalcium phosphate.
- 8. A method for inducing bone formation in a patient in need of same comprising administering to said patient an effective amount of a composition of Claim 1-7.
- 9. A process for producing BMP-1 comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-1, said DNA sequence being in relative association with an expression control sequence therefor, and isolating BMP-1 from said culture medium.
- 10. A process according to Claim 9 wherein said DNA sequence comprises substantially the nucleotide sequence of Table VI.
- 11. A process for producing BMP-2 Class I comprising culturing in a suitable culture medium a cell line transformed

with a DNA sequence encoding BMP-2 Class I, said DNA sequence being in relative association with an expression control sequence therefor, and isolating BMP-2 Class I from said culture medium.

- 12. A process for according to Claim 11 wherein said DNA sequence comprises substantially the nucleotide sequence of Table VII.
- 13. A process for producing BMP-2 Class II comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-2 Class II, said DNA sequence being in relative association with an expression control sequence therefor, and isolating BMP-2 Class II from said culture medium.
- 14. A process according to Claim 13 wherein said DNA sequence comprises substantially the nucleotide sequence of Table VIII.
- 15. A process for producing BMP-3 comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-3, said DNA sequence being in relative association with an expression control sequence therefor and isolating BMP-3 from said culture medium.
- 16. A process according to Claim 15 wherein said DNA sequence comprises substantially the nucleotide sequence of Table IVA and IVB.
- 17. A cDNA sequence encoding BMP-1 comprising substantially the nucleotide sequence of Table VI or a sequence which hybridize thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-1.
- 18. A cDNA sequence encoding BMP-2 Class I comprising substantially the nucleotide sequence of Table VII or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class I.
- 19. A cDNA sequence encoding BMP-2 Class II comprising

substantially the nucleotide sequence of Table VIII or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class II.

20. A cDNA sequence encoding BMP-3 comprising substantially the nucleotide sequence of Table IVA and IVB or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-3.

AMENDED CLAIMS

[received by the International Bureau on 8 December 1987 (08.12.87) original claims 6, 8, 10, 12, 14, 16-20 amended; new claims 21-23 added; other claims unchanged (13 pages)]

- 1. A pharmaceutical composition comprising a protein selected from the group consisting of:
 - (a) BMP-1;
 - (b) BMP-2 Class I;
 - (c) BMP-2 Class II;
 - (d) BMP-3; and

mixtures thereof, in a pharmaceutically acceptable vehicle.

- 2.. A composition of Claim 1 wherein said protein is BMP-1.
- 3. A composition of Claim 1 wherein said protein is BMP-2 Class I.
- 4. A composition of Claim 1 wherein said protein is BMP-2 Class II.
- 5. A composition of Claim 1 wherein said protein is BMP-3.
- 6. The pharmaceutical composition of Claim 1 further comprising a matrix capable of delivering the composition to the site of the bone or cartilage defect and providing a structure for inducing bone or cartilage formation.
- 7. The composition of Claim 6 wherein said matrix comprises a material selected from the group consisting of hydroxyapatite, collagen, polylactic acid and tricalcium phosphate.
- 8. A method for inducing bone or cartilage formation in a patient in need of same comprising administering to said patient an effective amount of a composition of Claim 1-7.

- 9. A process for producing BMP-1 comprising the steps of culturing in a suitable culture medium a host cell transformed with a DNA sequence encoding BMP-1, said DNA sequence being in relative association with an expression control sequence therefor; and isolating said BMP-1 therefrom.
- 10. A process according to Claim 9 wherein said DNA sequence comprises substantially the nucleotide sequence as follows:

10 20 30 50
CTAGAGGCCC CTTCCCTCGC CGCCCCCCG CCAGC ATG CCC GGC GTG GCC CTG CCG
MET Pro Gly Val Ala Arg Leu Pro

65 80 95 110
CTG CTG CTC GGG CTG CTG CTC CCG CGT CCC GGC CGG CCG CTG GAC TTG GCC
Leu Leu Gly Leu Leu Leu Leu Pro Arg Pro Gly Arg Pro Leu Asp Leu Ala

125 140 155 GAC TAC ACC TAT GAC CTG GCG GAG GAG GAC GAC TCG GAG CCC CTC AAC TAC AAA Asp Tyr Thr Tyr Asp Leu Ala Glu Glu Asp Asp Ser Glu Pro Leu Asn Tyr Lys

170 185 200 215
GAC CCC TGC AAG GCG GCT GCC TTT CTT GGG GAC ATT GCC CTG GAC GAA GAG GAC
Asp Pro Cys Lys Ala Ala Ala Phe Leu Gly Asp Ile Ala Leu Asp Glu Glu Asp

230 245 260 275 CTG AGG GCC TTC CAG GTA CAG CAG GCT GTG GAT CTC AGA CGG CAC ACA GCT CGT Leu Arg Ala Phe Gln Val Gln Gln Ala Val Asp Leu Arg Arg His Thr Ala Arg

290 305 320

AAG TOC TOC ATC AAA GCT GCA GIT COA GGA AAC ACT TOT ACC COC AGC TGC CAG Lys Ser Ser Ile Lys Ala Ala Val Pro Gly Asn Thr Ser Thr Pro Ser Cys Gln

335 350 365 380

AGC ACC AAC GGG CAG CCT CAG AGG GGA GCC TGT GGG AGA TGG AGA GGT AGA TCC

Ser Thr Asn Gly Gln Pro Gln Arg Gly Ala Cys Gly Arg Trp Arg Gly Arg Ser

395 410 425

OGT AGC CGG CGG GCG ACG TCC CGA CCA GAG CGT GTG TCG CCC GAT GGG GTC Arg Ser Arg Arg Ala Ala Thr Ser Arg Pro Glu Arg Val Trp Pro Asp Gly Val

440 455 470 485 ATC CCC TIT GTC ATT GGG GGA AAC TTC ACT GGT AGC CAG AGG GCA GTC TTC CGG Ile Pro Phe Val Ile Gly Gly Asn Phe Thr Gly Ser Gln Arg Ala Val Phe Arg

500 515 530 545 CAG GCC ATG AGG CAC TGG GAG AAG CAC ACC TGT GTC ACC TTC CTG GAG CGC ACT Gln Ala MET Arg His Trp Glu Lys His Thr Cys Val Thr Phe Leu Glu Arg Thr

590 575 560 GAC GAG GAC AGC TAT ATT GTG TTC ACC TAT OGA CCT TGC GGG TGC TGC TCC TAC Asp Glu Asp Ser Tyr Ile Val Phe Thr Tyr Arg Pro Cys Gly Cys Cys Ser Tyr 650 635 605 620 GTG GGT CGC CGC GGG GGC CCC CAG GCC ATC TCC ATC GGC AAG AAC TGT GAC Val Gly Arg Arg Gly Gly Pro Gln Ala Ile Ser Ile Gly Lys Asn Cys Asp 665 AAG TTC GGC ATT GTG GTC CAC GAG CTG GGC CAC GTC GTC GGC TTC TGG CAC GAA Lys Phe Gly Ile Val Val His Glu Leu Gly His Val Val Gly Phe Trp His Glu 740 CAC ACT CCG CCA GAC CCG GAC CCC CAC GIT TCC ATC GIT CGT GAG AAC ATC CAG His Thr Arg Pro Asp Arg Asp Arg His Val Ser Ile Val Arg Glu Asn Ile Gln CCA GGG CAG GAG TAT AAC TTC CTG AAG ATG GAG CCT CAG GAG GTG GAG TCC CTG Pro Gly Gln Glu Tyr Asn Phe Leu Lys MET Glu Pro Gln Glu Val Glu Ser Leu GGG GAG ACC TAT GAC TTC GAC AGC ATC ATG CAT TAC GCT CGG AAC ACA TTC TCC Gly Glu Thr Tyr Asp Phe Asp Ser Ile MET His Tyr Ala Arg Asn Thr Phe Ser 905 890 AGG GGC ATC TTC CTG GAT ACC ATT GTC CCC AAG TAT GAG GTG AAC GGG GTG AAA Arg Gly Ile Phe Leu Asp Thr Ile Val Pro Lys Tyr Glu Val Asn Gly Val Lys 935 CCT CCC ATT GGC CAA AGG ACA CGG CTC AGC AAG GGG GAC ATT GCC CAA GCC CGC Pro Pro Ile Gly Gln Arg Thr Arg Leu Ser Lys Gly Asp Ile Ala Gln Ala Arg 995 1010 AAG CTT TAC AAG TGC CCA GCC TGT GGA GAG ACC CTG CAA GAC AGC ACA GGC AAC Lys Leu Tyr Lys Cys Pro Ala Cys Gly Glu Thr Leu Gln Asp Ser Thr Gly Asn 1070 1055 1040 TTC TCC TCC CCT GAA TAC CCC AAT GGC TAC TCT GCT CAC ATG CAC TGC GTG TGG Phe Ser Ser Pro Glu Tyr Pro Asn Gly Tyr Ser Ala His MET His Cys Val Trp 1115 1130 1100 OGC ATC TCT GTC ACA CCC GGG GAG AAG ATC ATC CTG AAC TTC ACG TCC CTG GAC Arg Ile Ser Val Thr Pro Gly Glu Lys Ile Ile Leu Asn Phe Thr Ser Leu Asp 1190 1160 1175 1145 CTG TAC OGC AGC OGC CTG TGC TGG TAC GAC TAT GTG GAG GTC OGA GAT GGC TTC Leu Tyr Arg Ser Arg Leu Cys Trp Tyr Asp Tyr Val Glu Val Arg Asp Gly Phe 1220 1235 TGG AGG AAG GOG CCC CTC CGA GGC CGC TTC TGC GGG TCC AAA CTC CCT GAG CCT Trp Arg Lys Ala Pro Leu Arg Gly Arg Phe Cys Gly Ser Lys Leu Pro Glu Pro

1250 1265 1280 1295 ATC GIC TCC ACT GAC AGC CGC CTC TGG GIT GAA TIC CGC AGC AGC AGC AAT TGG Ile Val Ser Thr Asp Ser Arg Leu Trp Val Glu Phe Arg Ser Ser Ser Asn Trp 1310. 1325 1340 1355 GIT GGA AAG: GGC TIC TIT GCA GTC TAC GAA GCC ATC TGC GGG GGT GAT GTG AAA Val Gly Lys: Gly Phe Phe Ala Val Tyr Glu Ala Ile Cys Gly Gly Asp Val Lys AAG GAC TAT GGC CAC ATT CAA TOG COC AAC TAC CCA GAC GAT TAC CGG COC AGC Lys Asp Tyr Gly His Ile Gln Ser Pro Asn Tyr Pro Asp Asp Tyr Arg Pro Ser AAA GIC TGC ATC TGG CGG ATC CAG GIG TCT GAG GGC TIC CAC GIG GGC CTC ACA Lys Val Cys: Ile Trp Arq Ile Gln Val Ser Glu Gly Phe His Val Gly Leu Thr 1475 1490 1505 TTC CAG TCC TTT GAG ATT GAG CGC CAC GAC AGC TGT GCC TAC GAC TAT CTG GAG Phe Gln Ser Phe Glu Ile Glu Arg His Asp Ser Cys Ala Tyr Asp Tyr Leu Glu 1535 1550 GTG OGC GAC GGG CAC AGT GAG AGC AGC CTC ATC GGG CGC TAC TGT GGC TAT Val Arg Asp GTy His Ser Glu Ser Ser Thr Leu Ile Gly Arg Tyr Cys Gly Tyr 1595 1610 1625 CAG AAG CCT GAT GAC ATC AAG AGC ACG TCC AGC CGC CTC TGG CTC AAG TTC GTC Glu Lys Pro Asp Asp Ile Lys Ser Thr Ser Ser Arg Leu Tro Leu Lys Phe Val 1640 1655 TCT GAC GGG TCC ATT AAC AAA GCG GGC TIT GCC GTC AAC TIT TIC AAA GAG GTG Ser Asp Gly Ser Ile Asn Lys Ala Gly Phe Ala Val Asn Phe Phe Lys Glu Val 1685 1700 1715 1730 CAC GAG TGC TCT CGG CCC AAC CGC GGG GGC TGT GAG CAG CGG TGC CTC AAC ACC Asp Glu Cys Ser Arg Pro Asn Arg Gly Gly Cys Glu Gln Arg Cys Leu Asn Thr 1745 17.60 1775 CTG GGC AGC TAC AAG TGC AGC TGT GAC CCC GGG TAC GAG CTG GCC CCA GAC AAG Leu Gly Ser Tyr Lys Cys Ser Cys Asp Pro Gly Tyr Glu Leu Ala Pro Asp Lys 1790 1805 1820 OGC OGC TGT GAG GCT GCT TGT GGC GGA TTC CTC ACC AAG CTC AAC GGC TCC ATC Arg Arg Cys Glu Ala Ala Cys Gly Gly Phe Leu Thr Lys Leu Asn Gly Ser Ile 1865 ACC AGC COG GGC TGG CCC AAG GAG TAC CCC CCC AAC AAG AAC TGC ATC TGG CAG Thr Ser Pro Gly Trp Pro Lys Glu Tyr Pro Pro Asn Lys Asn Cys Ile Trp Gln 1910 1925 CTG GTG GCC CCC ACC CAG TAC CGC ATC TCC CTG CAG TTT GAC TTC TTT GAG ACA Leu Val Ala Pro Thr Gln Tyr Arg Ile Ser Leu Gln Phe Asp Phe Phe Glu Thr

1955 1970 1985 2000
GAG GGC AAT GAT GTG TGC AAG TAC GAC TTC GTG GAG GTG CGC AGT GGA CTC ACA
Glu Gly Asn Asp Val Cys Lys Tyr Asp Phe Val Glu Val Arg Ser Gly Leu Thr

2015 2030 2045
GCT GAC TCC AAG CTG CAT GGC AAG TTC TGT GGT TCT GAG AAG CCC GAG GTC ATC
Ala Asp Ser Lys Leu His Gly Lys Phe Cys Gly Ser Glu Lys Pro Glu Val Ile

2060 2075 2090 2105

ACC TCC CAG TAC AAC AAC ATG CGC GTG GAG TTC AAG TCC GAC AAC ACC GTG TCC

Thr Ser Gln Tyr Asn Asn MET Arg Val Glu Phe Lys Ser Asp Asn Thr Val Ser

2120 2135 2150 2165

AAA AAG GGC TTC AAG GCC CAC TTC TTC TCA GAA AAG AGG CCA GCT CTG CAG CCC
Lys Lys Gly Phe Lys Ala His Phe Phe Ser Glu Lys Arg Pro Ala Leu Gln Pro

2180 2195 2210
CCT CGG GGA CGC CCC CAC CAG CTC AAA TTC CGA GTG CAG AAA AGA AAC CGG ACC
Pro Arg Gly Arg Pro His Gln Leu Lys Phe Arg Val Gln Lys Arg Asn Arg Thr

2225 2235 2245 2255 2265 2275 2285 CCC CAG TGAGGCCTGC CAGGCCTCCC GGACCCCTTG TTACTCAGGA ACCTCACCTT GGACGGAATG Pro Gln

2295 2305 2315 2325 2335 2345 2355 GGATGGGGGC TTGGGTGCCC ACCAACCCCC CACCTCCACT CTGCCATTCC GGCCCACCTC CCTCTGGCCG

2365 2375 2385 2395 2405 2415 2425 GACAGAACTG GTGCTCTCTT CTCCCCACTG TGCCCGTCCG CGGACCGGGG ACCCTTCCCC GTGCCCTACC

2435 2445 2455 2465 2475 2485 2495 CCCTCCCATT TTGATGGTGT CTGTGACATT TCCTGTTGTG AAGTAAAAGA GGGACCCCTG CGTCCTGCCT

CTAGA

- 11. A process for producing BMP-2 Class I comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-2 Class I, said DNA sequence being in relative association with an expression control sequence therefor, and isolating BMP-2 Class I from said culture medium.
- 12. A process for according to Claim 11 wherein said DNA sequence comprises substantially the nucleotide sequence as

follows:

10 20 30 40 50 60 70 GIOGACICIA GAGIGIGIGI CAGCACTIGG CIGGGGACIT CTIGAACTIG CAGGGAGAAT AACITGCGCA
80 90 100 110 120 130 140 CCCCACTITG CGCGGGCCCAG CGCAGCCTGC TTCGCCATCT CCGAGCCCCA CCGCCCCTCC
150 150 170 180 190 200 210 ACTCCTOGGC CITGCCCGGC ACTCAGACGC TGITCCCAGC GTGAAAAGAG AGACTGCGGG GCCGCCACCC
220 230 240 250 260 270 280 GGGAGAAGGA GGAGGAAAG AAAAGGAACG GACATTCGGT CCTTGCGCCA GGTCCTTTGA CCAGAGTTTT
290 300 310 320 330 340 350 TCCATGIGGA CGCICITICA ATGGACGIGI CCCCGCGIGC TICITAGACG GACIGCGGIC TCCIAAAGGI
370 385 400 CGACCIATG GIG GCC GGG ACC CGC TGT CIT CTA GCG TIG CIT CCC CAG GIC MET Val Ala Gly Thr Arg Cys Leu Leu Ala Leu Leu Pro Gln Val
415 430 445 CTC CTG:GGC:GGC GCC GCC CTC GTT CCG GAG CTG GGC CGC AGG AAG TTC GCG Leu Leu Gly Gly Ala Ala Gly Leu Val Pro Glu Leu Gly Arg Arg Lys Phe Ala
460 475 490 505 GOG GOG TOG TOG GGC CGC CCC TCA TCC CAG CCC TCT GAC GAG GTC CTG AGC GAG Ala Ala Ser Ser Gly Arg Pro Ser Ser Gln Pro Ser Asp Glu Val Leu Ser Glu
520 535 550 565 TTC GAG TTG CGG CTG AGC ATG TTC GGC CTG AAA CAG AGA CCC ACC CCC AGC Phe Glu Leu Arg Leu Leu Ser MET Phe Gly Leu Lys Gln Arg Pro Thr Pro Ser
580 595 610 AGG GAC GCC GTG GTG CCC CCC TAC ATG CTA GAC CTG TAT CGC AGG CAC TCG GGT Arg Asp Ala Val Val Pro Pro Tyr MET Leu Asp Leu Tyr Arg Arg His Ser Gly
625 640 655 670 CAG CCG GGC TCA CCC GCC CCA CAC CAC CGG TTG GAG AGG GCA GCC AGC CGA GCC Gln Pro Gly Ser Pro Ala Pro Asp His Arg Leu Glu Arg Ala Ala Ser Arg Ala
685 700 715 AAC ACT GTG CGC AGC TTC CAC CAT GAA GAA TCT TTG GAA GAA CTA CCA GAA ACG Asn Thr Val Arg Ser Phe His His Glu Glu Ser Leu Glu Glu Leu Pro Glu Thr
730 745 760 775 AGT GGG AAA ACA ACC CGG AGA TTC TTC TTT AAT TTA AGT TCT ATC CCC ACG GAG Ser Gly Lys Thr Thr Arg Arg Phe Phe Phe Asn Leu Ser Ser Ile Pro Thr Glu
790 805 820 835 GAG TIT ATC ACC TCA GCA GAG CIT CAG GIT TIC CGA GAA CAG ATG CAA GAT GCT Glu Phe Ile Thr Ser Ala Glu Leu Gln Val Phe Arg Glu Gln MET Gln Asp Ala

865 880 850 TTA GGA AAC AAT AGC AGT TTC CAT CAC CGA ATT AAT ATT TAT GAA ATC ATA AAA Leu Gly Asn Asn Ser Ser Phe His His Arg Ile Asn Ile Tyr Glu Ile Ile Lys 925 910 CCT GCA ACA GCC AAC TCG AAA TTC CCC GTG ACC AGT CTT TTG GAC ACC AGG TTG Pro Ala Thr Ala Asn Ser Lys Phe Pro Val Thr Ser Leu Leu Asp Thr Arg Leu 970 GTG AAT CAG AAT GCA AGC AGG TGG GAA AGT TIT GAT GTC ACC CCC GCT GTG ATG Val Asn Gln Asn Ala Ser Arg Trp Glu Ser Phe Asp Val Thr Pro Ala Val MET 1030 1000 1015 CGG TGG ACT GCA CAG GGA CAC GCC AAC CAT GGA TTC GTG GTG GAA GTG GCC CAC Arg Trp Thr Ala Gln Gly His Ala Asn His Gly Phe Val Val Glu Val Ala His 1090 1060 1075 TIG GAG GAG AAA CAA GGI GTC TCC AAG AGA CAT GIT AGG ATA AGC AGG TCT TIG Leu Glu Glu Lys Gln Gly Val Ser Lys Arg His Val Arg Ile Ser Arg Ser Leu 1135 1120 CAC CAA GAT GAA CAC AGC TGG TCA CAG ATA AGG CCA TTG CTA GTA ACT TTT GGC His Gln Asp Glu His Ser Trp Ser Gln Ile Arg Pro Leu Leu Val Thr Phe Gly 1195 1210 1165 1180 CAT GAT GGA AAA GGG CAT CCT CTC CAC AAA AGA GAA AAA CGT CAA GCC AAA CAC His Asp Gly Lys Gly His Pro Leu His Lys Arg Glu Lys Arg Gln Ala Lys His 1225 1240 1255 AAA CAG CGG AAA CGC CIT AAG TCC AGC TGT AAG AGA CAC CCT TTG TAC GTG GAC Lys Gln Arg Lys Arg Leu Lys Ser Ser Cys Lys Arg His Pro Leu Tyr Val Asp 1270 1285 1300 TIC AGT GAC GIG GGG TGG AAT GAC TGG AIT GIG GCT CCC CCG GGG TAT CAC GCC Phe Ser Asp Val Gly Trp Asn Asp Trp Ile Val Ala Pro Pro Gly Tyr His Ala 1345 1330 TTT TAC TGC CAC GGA GAA TGC CCT TTT CCT CTG GCT GAT CAT CTG AAC TCC ACT Phe Tyr Cys His Gly Glu Cys Pro Phe Pro Leu Ala Asp His Leu Asn Ser Thr 1405 AAT CAT GCC ATT GTT CAG ACG TTG GTC AAC TCT GTT AAC TCT AAG ATT CCT AAG Asn His Ala Ile Val Gln Thr Leu Val Asn Ser Val Asn Ser Lys Ile Pro Lys 1450 1465 GCA TGC TGT GTC CCG ACA GAA CTC AGT GCT ATC TCG ATG CTG TAC CTT GAC GAG Ala Cys Cys Val Pro Thr Glu Leu Ser Ala Ile Ser MET Leu Tyr Leu Asp Glu

1495 1510 1525

AAT GAA AAG GIT GTA TTA AAG AAC TAT CAG GAC ATG GIT GIG GAG GGT TGT GGG

Asn Glu Lys Val Val Leu Lys Asn Tyr Gln Asp MET Val Val Glu Gly Cys Gly

1540 1553 1563 1573 1583 1593 1603 TGT CGC TAGTACAGCA AAATTAAATA CATAAATATA TATATATATA TATATTITTAG AAAAAAGAAA Cys Arg

AAAA

- 13. A process for producing BMP-2 Class II comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-2 Class II, said DNA sequence being in relative association with an expression control sequence therefor, and isolating BMP-2 Class II from said culture medium.
- 14. A process according to Claim 13 wherein said DNA sequence comprises substantially the nucleotide sequence as follows:

10 20 30 40 50 60 70 CTCTAGAGGG CAGAGGAGGA GGGAGGGAGG GAAGGAGGGC GGAGCCCGGC CCCGAAGCTA GGTGAGTGTG

80 90 100 110 120 130 140 GCATCOGAGC TGAGGGACCC GAGCCTGAGA CGCCCCTGCT GCTCCGGCTG AGTATCTAGC TTGTCTCCCC

220 230 240 250 260 270 280 CAACOGITCA GAGGICCOCA GGAGCIGCIG CICGOGAGCC CGCTACIGCA GGGACCIATG GAGCCATTCC

290 300 310 320 330 340 350 GTAGTGCCAT CCCGAGCAAC GCACTGCTGC AGCTTTCCTG AGCCTTTCCA GCAAGTTTGT TCAAGATTCG

360 370 380 390 400
CIGICAAGAA TCATGGACTG TTATTATATG CCTTGTTTTC TGTCAAGACA CC ATG ATT CCT
MET Ile Pro

417 432 447 462
GGT AAC CGA ATG CTG ATG GTC GTT TTA TTA TGC CAA GTC CTG CTA GGA GGC GCG
Gly Asn Arg MET Leu MET Val Val Leu Leu Cys Gln Val Leu Leu Gly Gly Ala

477 492 507 AGC CAT GCT AGT TIG ATA CCT GAG ACG GGG AAG AAA AAA GTC GCC GAG ATT CAG Ser His Ala Ser Leu Ile Pro Glu Thr Gly Lys Lys Lys Val Ala Glu Ile Gln

552 537 522 GGC CAC GOG GGA GGA CGC CGC TCA GGG CAG AGC CAT GAG CTC CTG CGG GAC TTC Gly His Ala Gly Gly Arg Arg Ser Gly Gln Ser His Glu Leu Leu Arg Asp Phe 612 597 CAG GOG ACA CIT CIG CAG ATG TIT GGG CIG CGC CGC CGC CAG CCT AGC AAG Glu Ala Thr Leu Leu Gln MET Phe Gly Leu Arg Arg Arg Pro Gln Pro Ser Lys 657 642 AGT GCC GTC ATT CCG GAC TAC ATG CGG GAT CTT TAC CGG CTT CAG TCT GGG GAG Ser Ala Val Ile Pro Asp Tyr MET Arg Asp Leu Tyr Arg Leu Gln Ser Gly Glu 717 687 702 GAG GAG GAA GAG CAG ATC CAC AGC ACT GGT CTT GAG TAT CCT GAG CGC CCC GCC Glu Glu Glu Glu Gln Ile His Ser Thr Gly Leu Glu Tyr Pro Glu Arg Pro Ala 762 747 AGC OGG GCC AAC ACC GTG AGG AGC TTC CAC CAC GAA GAA CAT CTG GAG AAC ATC Ser Arg Ala Asn Thr Val Arg Ser Phe His His Glu Glu His Leu Glu Asn Ile 792 807 822 CCA GGG ACC AGT GAA AAC TCT GCT TTT CGT TTC CTC TTT AAC CTC AGC AGC ATC Pro Gly Thr Ser Glu Asn Ser Ala Phe Arg Phe Leu Phe Asn Leu Ser Ser Ile 852 867 882 CCT GAG AAC GAG GCG ATC TCC TCT GCA GAG CTT CGG CTC TTC CGG GAG CAG GTG Pro Glu Asn Glu Ala Ile Ser Ser Ala Glu Leu Arg Leu Phe Arg Glu Gln Val 927 942 912 GAC CAG GGC CCT GAT TGG GAA AGG GGC TTC CAC CGT ATA AAC ATT TAT GAG GTT Asp Gln Gly Pro Asp Trp Glu Arg Gly Phe His Arg Ile Asn Ile Tyr Glu Val 987 957 972 ATG AAG CCC CCA GCA GAA GTG GTG CCT GGG CAC CTC ATC ACA CGA CTA CTG GAC MET Lys Pro Pro Ala Glu Val Val Pro Gly His Leu Ile Thr Arg Leu Leu Asp 1017 1032 ACG AGA CTG GTC CAC CAC AAT GTG ACA CGG TGG GAA ACT TTT GAT GTG AGC CCT Thr Arg Leu Val His His Asn Val Thr Arg Trp Glu Thr Phe Asp Val Ser Pro 1092 1077 GOG GTC CTT CGC TGG ACC CGG GAG AAG CAG CCA AAC TAT GGG CTA GCC ATT GAG Ala Val Leu Arg Trp Thr Arg Glu Lys Gln Pro Asn Tyr Gly Leu Ala Ile Glu 1152 1137 GTG ACT CAC CTC CAT CAG ACT CGG ACC CAC CAG GGC CAG CAT GTC AGG ATT AGC Val Thr His Leu His Gln Thr Arg Thr His Gln Gly Gln His Val Arg Ile Ser 1197 1182 CGA TOG TTA CCT CAA GGG AGT GGG AAT TGG GCC CAG CTC CGG CCC CTC CTG GTC Arg Ser Leu Pro Gln Gly Ser Gly Asn Trp Ala Gln Leu Arg Pro Leu Leu Val

CGGAATTC

ACC TIT GGC CAT GAT GGC CGG GGC CAT GCC TIG ACC CGA CGC CGG AGG GCC AAG Thr Phe Gly His Asp Gly Arg Gly His Ala Leu Thr Arg Arg Arg Arg Ala Lys OFF AGC OCT AAG CAT CAC TCA CAG OGG GCC AGG AAG AAG AAT AAG AAC TGC OGG Arg Ser Pro Lys His His Ser Gln Arg Ala Arg Lys Lys Asn Lys Asn Cys Arg OGC CAC TOG CIC TAT GIG GAC TIC AGC GAT GIG GGC TGG AAT GAC TGG AIT GIG Arg His Ser Leu Tyr Val Asp Phe Ser Asp Val Gly Trp Asn Asp Trp Ile Val GCC CCA CCA GGC TAC CAG GCC TTC TAC TGC CAT GGG GAC TGC CCC TTT CCA CTG Ala Pro Pro Gly Tyr Gln Ala Phe Tyr Cys His Gly Asp Cys Pro Phe Pro Leu GCT GAC CAC CTC AAC TCA ACC AAC CAT GCC ATT GTG CAG ACC CTG GTC AAT TCT Ala Asp His Leu Asn Ser Thr Asn His Ala Ile Val Gln Thr Leu Val Asn Ser GTC AAT TOO AGI ATO COO AAA GOO TGI TGI GIG COO ACI GAA CIG AGI GOO ATO Val Asn Ser Ser Ile Pro Lys Ala Cys Cys Val Pro Thr Glu Leu Ser Ala Ile TCC ATG CTG TAC CTG GAT GAG TAT GAT AAG GTG GTA CTG AAA AAT TAT CAG GAG Ser MET Leu Tyr Leu Asp Glu Tyr Asp Lys Val Val Leu Lys Asn Tyr Gln Glu ATG GTA GTA GAG GGA TGT GGG TGC CGC TGAGATCAGG CAGTCCTTGA GGATAGACAG ATATACACAC MET Val Val Glu Gly Cys Gly Cys Arg CACACACACA CACCACATAC ACCACACACA CACGITCCCA TCCACTCACC CACACACTAC ACACACTGCT TCCTTATAGC TGGACTTTTA TTTAAAAAAA AAAAAAAAA AATGGAAAAA ATCCCTAAAC ATTCACCTTG ACCITATITA TGACIPTACG TGCAAATGIT TIGACCATAT TGATCATATA TITTGACAAA ATATATITAT AACTACGTAT TAAAAGAAAA AAATAAAATG AGTCATTATT TTAAAAAAAA AAAAAAACT CTAGAGTCGA

15. A process for producing BMP-3 comprising culturing in a suitable culture medium a cell line transformed with a DNA sequence encoding BMP-3, said DNA sequence being in relative

association with an expression control sequence therefor and isolating BMP-3 from said culture medium.

16. A process according to Claim 15 wherein said DNA sequence comprises substantially the nucleotide sequence as follows:

383 393 403 413 428

GAGGAGGAAG CCCTCTACCG GCGTCCTTCT GCCTCTGCAG AAC AAT GAG CTT CCT GGG GCA
Asn Asn Glu Leu Pro Gly Ala

443 458 473 488
GAA TAT CAG TAC AAG GAG GAT GAA GTA TGG GAG GAG AGG AAG CCT TAC AAG ACT
Glu Tyr Gln Tyr Lys Glu Asp Glu Val Trp Glu Glu Arg Lys Pro Tyr Lys Thr

503 518 533
CTT CAG ACT CAG CCC CCT GAT AAG AGT AAG AAC AAA AAG AAA CAG AGG AAG GGA
Leu Gln Thr Gln Pro Pro Asp Lys Ser Lys Asn Lys Lys Lys Gln Arg Lys Gly

548 563 578 593
CCT CAG CAG AAG AGT CAG ACG CTC CAG TTT GAT GAA CAG ACC CTG AAG AAG GCA
Pro Gln Gln Lys Ser Gln Thr Leu Gln Phe Asp Glu Gln Thr Leu Lys Lys Ala

AGA AGA AAG CAA TGG ATT GAA CCC CGG AAT TGT GCC AGA CGG TAC CTT AAA GTG Arg Arg Lys Gln Trp Ile Glu Pro Arg Asn Cys Ala Arg Arg Tyr Leu Lys Val

GAC TTC GCA GAT ATT GGC TGG AGC GAA TGG ATT ATT TCC CCC AAG TCC TTC GAT Asp Phe Ala Asp Ile Gly Trp Ser Glu Trp Ile Ile Ser Pro Lys Ser Phe Asp

713 728 743 756 766
GCC TAT TAC TGC TCC GGA GCG TGC CAG TTC CCC ATG CCA AAG GTAGCCATTG TTTTTTGTCC
Ala Tyr Tyr Cys Ser Gly Ala Cys Gln Phe Pro MET Pro Lys

776 786
TGICCITCCC ATTICCATAG; and

284 294 304 319
CTAACCTGTG TTCTCCCTTT TCGTTCCTAG TCT TTG AAG CCA TCA AAT CAC GCT ACC
Ser Leu Lys Pro Ser Asn His Ala Thr

334 349 364 379
ATC CAG AGT ATA GTG AGA GCT GTG GGG GTC GTC CCT GGA ATC CCC GAG CCT TGC
Ile Gln Ser Ile Val Arg Ala Val Gly Val Val Pro Gly Ile Pro Glu Pro Cys

394 409 . 424 439
TGT GTG CCA GAA AAG ATG TCC TCA CTC AGC ATC TTA TTC TTT GAT GAA AAC AAG
Cys Val Pro Glu Lys MET Ser Ser Leu Ser Ile Leu Phe Phe Asp Glu Asn Lys

454 469 484

AAT GIG GTA CIT AAA GTA TAT CCA AAC ATG ACA GTA GAG TCT TGT GCT TGC AGA
Asn Val Val Leu Lys Val Tyr Pro Asn MET Thr Val Glu Ser Cys Ala Cys Arg

503 513 523 533 TAACCTGGIG AAGAACTCAT CIGGATGCIT AACTCAATCG.

- 17. A cDNA sequence encoding BMP-1 comprising substantially the nucleotide sequence recited in Claim 10 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-1.
- 18. A cDNA sequence encoding BMP-2 Class I comprising substantially the nucleotide sequence recited in Claim 12 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class I.
- 19. A cDNA sequence encoding BMP-2 Class II comprising substantially the nucleotide sequence recited in Claim 14 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class II.
- 20. A cDNA sequence encoding BMP-3 comprising substantially the nucleotide sequence recited in Claim 16 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-3.
- 21. A vector containing a DNA sequence encoding an osteoinductive protein and heterologous DNA, the DNA sequence encoding the protein being selected from the group consisting of:
- a. a DNA sequence encoding BMP-1 comprising substantially the nucleotide sequence recited in Claim 10 or a sequence which

hybridize thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-1;

- b. a DNA sequence encoding BMP-2 Class I comprising substantially the nucleotide sequence recited in Claim 12 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class I;
- c. a DNA sequence encoding BMP-2 Class II comprising substantially the nucleotide sequence recited in Claim 14 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-2 Class II; and
- d. a DNA sequence encoding BMP-3 comprising substantially the nucleotide sequence recited in Claim 16 or a sequence which hybridizes thereto under stringent conditions and which upon expression codes for a protein exhibiting substantial properties of BMP-3.
- 22. A cell transformed with a vector according to claim 21 which is capable of expressing a DNA sequence encoding the osteoinductive protein and progeny of said cell.
- 23. The transformed cell according to claim 24 selected from the group consisting of a mammalian cell, a bacterial cell, an insect cell, and a yeast cell.

INTERNATIONAL SEARCH REPORT

International Application No PCT/US87/01537

I. CLASS	IFICATION OF SUBJECT MATTER (if several class	sification symbols apply, indicate all) 3	1/000//0133/_
According	to International Patent Classification (IPC) or to both Na): C07K 13/00,15/00; A61K : 530/350,395,397; 514/12	itional Classification and IPC 37/00; See Attachme	nt hment
	S SEARCHED	, , , , , , , , , , , , , , , , , , , ,	
	. Minimum Docume	entation Searched +	
Classification	on System	Classification Symbols	
ບຣ	530/350,395,397; 514 435/68, 70, 172.3; 9		
	Documentation Searched other to the Extent that such Document	than Minimum Documentation s are Included in the Fields Searched 5	
COMPU'	TER SEARCH CAS, APS: BONE I TIVE PROTEIN, BMP, OSTEOIN	MORPHOGEN, BONE DUCTIVE FACTOR	
III. DOCU	MENTS CONSIDERED TO BE RELEVANT !+		
Category •-	Citation of Document, 16 with indication, where ap		Relevant to Claim No. 1
$\frac{X,P}{Y,P}$	US, A, 4,619,989 (URIST	r) 28 Oct 1986.	$\frac{1-8}{9-20}$
$\frac{\mathbf{X}}{\mathbf{Y}}$	US, A, 4,563,350 (NATH) 7 January 1986.	AN ET AL)	1-8 9-20
$\frac{X}{Y}$	US, A, 4,455,256 (URIST	r) 19 June 1984.	1-8 9-20
x	Proc. Natl. Acad. Sci Uissued January 1984, (VD.C.),	JSA, Vol. 81,	1
Y	(URIST), "Purification morphogenetic protein be chromatography", pages	y hydroxyapatite	2-20
			1
		,	
"A" docu cons "E" earli	categones of cited documents: 13 ment defining the general state of the art which is not lidered to be of particular relevance or document but published on or after the international	"T" later document published after or priority date and not in conflicited to understand the princip invention "X" document of particular relevants	lict with the application but le or theory underlying the ace; the claimed invention
"L" docu whic citati "O" docu	idate iment which may throw doubts on priority claim(s) or h is cited to establish the publication date of another ion or other special reason (as specified) iment referring to an oral disclosure, use, exhibition or r means	cannot be considered novel of involve an inventive step "Y" document of particular relevant cannot be considered to involve document is combined with one ments, such combination being	r cannot be considered to nce; the claimed invention an inventive step when the or more other such docu-
"P" docu	ment published prior to the international filing date but than the priority date claimed	in the art. "&" document member of the same	
IV. CERTI	FICATION		
	Actual Completion of the International Search :	Date of Mailing of this International S	earch Report ²
	08 October 1987	Signature of Authorized Officer *0	11
	ISA/US	Alvin E. Tanenholtz	<u> </u>
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PCT/US87/01537

Attachment To Form PCT/ISA/210, Part I.

IPC(4): C12P 21/00, 21/02; C12N 15/00; C07H 15/12

US CL: 435/68, 70, 172.3; 935/13

III. DOCUM	III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)							
Category *	Citation of Document, 16 with indication, where appropriate, of the relevant passages 17	Relevant to Claim No 18						
Y	Science, Vol. 220 issued 13 May 1983 (Washington, D.C.) (URIST) "Bone cell Differentiation and Growth Factors" pages 680-686.	1-20						
Y	Proc. Natl. Acad Sci, USA, Vol. 80 issued November 1983 (Washington, D.C.) (SAMPATH ET AL), "Homology of bone-inductive proteins from human monkey, bovine and rat extracellular matrix," pages 6591-6595.	1-20						
Y	Proc. Natl. Acad. Sci, USA, Vol. 78 issued November 1981, (Washington, D.C.) (SUGGS ET AL), "Use of synthetic oligonucleotides as hybridization probes: Isolation of cloned cDNA sequence for human \$2-microglobulin" pages 6613-6617.	1-20						